

Tue Jul 17 13:24:47 2001 [BLASTP 2.1.3 [Apr-1-2001], NCBI]
/home/ruby/va/Molbio/carpenda/temp1/p1.DNA37140 (1119 aa)

Sequences pr 1 P_AAB481 2 P_AAB802 3 P_AAY706 4 P_AAY081 5 P_AAY133 6 P_AAB802 7 P_AAY706 8 P_AAY133 9 P_AAY080	62 H 63 H 74 H 14 H 95 H 61 H 72 H 93 H	ing High-scoring Segment Pairs:  Human PRO326 polypeptide - Homo sapiens.  Human PRO326 protein - Homo sapiens.  Protein PRO326 - Homo sapiens.  Human PRO335 protein - Homo sapiens.  Human PRO335 protein - Homo sapiens.  Protein PRO335 - Homo sapiens.  Human PRO335 protein - Homo sapiens.  Protein PRO335 protein - Homo sapiens.  Human PRO335 protein - Homo sapiens.  Human PRO335 protein - Homo sapiens.  Score Match Pct E-val  5870 1119 100 0.0  5
Score = 587	0 (22	man PRO326 polypeptide - Homo sapiens. (1119 aa) [1 seg] 265 bits), Expect = 0.0 19/1119 (100%), Positives = 1119/1119 (100%), at 1,1-1119,1119
DNA37140	1	MSAPSLRARAAGLGLLLCAVLGRAGRSDSGGRGELGQPSGVAAERPCPTTCRCLGDLLDC
P_AAB48162	1	**************************************
DNA37140	61	SRKRLARLPEPLPSWVARLDLSHNRLSFIKASSMSHLQSLREVKLNNNELETIPNLGPVS
P_AAB48162	61	**************************************
DNA37140	121	ANITLLSLAGNRIVEILPEHLKEFQSLETLDLSSNNISELQTAFPALQLKYLYLNSNRVT
P AAB48162	121	**************************************
- DNA37140	181	SMEPGYFDNLANTLLVLKLNRNRISAIPPKMFKLPQLQHLELNRNKIKNVDGLTFQGLGA
P AAB48162		**************************************
_		
DNA37140	241	LKSLKMQRNGVTKLMDGAFWGLSNMEILQLDHNNLTEITKGWLYGLLMLQELHLSQNAIN ***********************************
P_AAB48162	241	LKSLKMQRNGVTKLMDGAFWGLSNMEILQLDHNNLTEITKGWLYGLLMLQELHLSQNAIN
DNA37140	301	RISPDAWEFCQKLSELDLTFNHLSRLDDSSFLGLSLLNTLHIGNNRVSYIADCAFRGLSS
P_AAB48162	301	RISPDAWEFCQKLSELDLTFNHLSRLDDSSFLGLSLLNTLHIGNNRVSYIADCAFRGLSS
DNA37140	361	LKTLDLKNNEISWTIEDMNGAFSGLDKLRRLILQGNRIRSITKKAFTGLDALEHLDLSDN
P_AAB48162	361	LKTLDLKNNEISWTIEDMNGAFSGLDKLRRLILQGNRIRSITKKAFTGLDALEHLDLSDN
DNA37140	421	AIMSLQGNAFSQMKKLQQLHLNTSSLLCDCQLKWLPQWVAENNFQSFVNASCAHPQLLKG
P_AAB48162	421	**************************************
DNA37140	481	RSIFAVSPDGFVCDDFPKPQITVQPETQSAIKGSNLSFICSAASSSDSPMTFAWKKDNEL
P_AAB48162	481	**************************************
DNA37140	541	LHDAEMENYAHLRAQGGEVMEYTTILRLREVEFASEGKYQCVISNHFGSSYSVKAKLTVN

P_AAB48162	541	LHDAEMENYAHLRAQGGEVMEYTTILRLREVEFASEGKYQCVISNHFGSSYSVKAKLTVN
DNA37140	601	MLPSFTKTPMDLTIRAGAMARLECAAVGHPAPQIAWQKDGGTDFPAARERRMHVMPEDDV
P_AAB48162	601	MLPSFTKTPMDLTIRAGAMARLECAAVGHPAPQIAWQKDGGTDFPAARERRMHVMPEDDV
DNA37140	661	FFIVDVKIEDIGVYSCTAQNSAGSISANATLTVLETPSFLRPLLDRTVTKGETAVLQCIA
P_AAB48162	661	FFIVDVKIEDIGVYSCTAQNSAGSISANATLTVLETPSFLRPLLDRTVTKGETAVLQCIA
DNA37140	721	GGSPPPKLNWTKDDSPLVVTERHFFAAGNQLLIIVDSDVSDAGKYTCEMSNTLGTERGNV
P_AAB48162	721	GGSPPPKLNWTKDDSPLVVTERHFFAAGNQLLIIVDSDVSDAGKYTCEMSNTLGTERGNV
DNA37140	781	RLSVIPTPTCDSPQMTAPSLDDDGWATVGVVIIAVVCCVVGTSLVWVVIIYHTRRRNEDC
P_AAB48162	781	RLSVIPTPTCDSPQMTAPSLDDDGWATVGVVIIAVVCCVVGTSLVWVVIIYHTRRRNEDC
DNA37140	841	SITNTDETNLPADIPSYLSSQGTLADRQDGYVSSESGSHHQFVTSSGAGFFLPQHDSSGT
P_AAB48162	841	SITNTDETNLPADIPSYLSSQGTLADRQDGYVSSESGSHHQFVTSSGAGFFLPQHDSSGT
DNA37140	901	CHIDNSSEADVEAATDLFLCPFLGSTGPMYLKGNVYGSDPFETYHTGCSPDPRTVLMDHY
P_AAB48162	901	CHIDNSSEADVEAATDLFLCPFLGSTGPMYLKGNVYGSDPFETYHTGCSPDPRTVLMDHY
DNA37140	961	EPSYIKKKECYPCSHPSEESCERSFSNISWPSHVRKLLNTSYSHNEGPGMKNLCLNKSSL
P_AAB48162	961	EPSYIKKKECYPCSHPSEESCERSFSNISWPSHVRKLLNTSYSHNEGPGMKNLCLNKSSL
DNA37140	1021	DFSANPEPASVASSNSFMGTFGKALRRPHLDAYSSFGQPSDCQPRAFYLKAHSSPDLDSG
P_AAB48162	1021	DFSANPEPASVASSNSFMGTFGKALRRPHLDAYSSFGQPSDCQPRAFYLKAHSSPDLDSG
DNA37140	1081	SEEDGKERTDFQEENHICTFKQTLENYRTPNFQSYDLDT
P_AAB48162	1081	SEEDGKERTDFQEENHICTFKQTLENYRTPNFQSYDLDT
Score = 58	70 (2	man PRO326 protein - Homo sapiens. (1119 aa) [1 seg] 265 bits), Expect = 0.0 19/1119 (100%), Positives = 1119/1119 (100%), at 1,1-1119,1119
DNA37140	1	MSAPSLRARAAGLGLLLCAVLGRAGRSDSGGRGELGQPSGVAAERPCPTTCRCLGDLLDC
P_AAB80263	1	MSAPSLRARAAGLGLLLCAVLGRAGRSDSGGRGELGQPSGVAAERPCPTTCRCLGDLLDC
DNA37140	61	SRKRLARLPEPLPSWVARLDLSHNRLSFIKASSMSHLQSLREVKLNNNELETIPNLGPVS ************************************
P_AAB80263	61	SRKRLARLPEPLPSWVARLDLSHNRLSFIKASSMSHLQSLREVKLNNNELETIPNLGPVS
DNA37140	121	ANITLLSLAGNRIVEILPEHLKEFQSLETLDLSSNNISELQTAFPALQLKYLYLNSNRVT
P_AAB80263	121	ANITLLSLAGNRIVEILPEHLKEFQSLETLDLSSNNISELQTAFPALQLKYLYLNSNRVT
DNA37140	181	SMEPGYFDNLANTLLVLKLNRNRISAIPPKMFKLPQLQHLELNRNKIKNVDGLTFQGLGA
P_AAB80263	181	SMEPGYFDNLANTLLVLKLNRNRISAIPPKMFKLPQLQHLELNRNKIKNVDGLTFQGLGA

DNA37140	241	LKSLKMQRNGVTKLMDGAFWGLSNMEILQLDHNNLTEITKGWLYGLLMLQELHLSQNAIN
P_AAB80263	241	LKSLKMQRNGVTKLMDGAFWGLSNMEILQLDHNNLTEITKGWLYGLLMLQELHLSQNAIN
DNA37140	301	RISPDAWEFCQKLSELDLTFNHLSRLDDSSFLGLSLLNTLHIGNNRVSYIADCAFRGLSS ***********************************
P_AAB80263	301	RISPDAWEFCQKLSELDLTFNHLSRLDDSSFLGLSLLNTLHIGNNRVSYIADCAFRGLSS
DNA37140	361	LKTLDLKNNEISWTIEDMNGAFSGLDKLRRLILQGNRIRSITKKAFTGLDALEHLDLSDN
P_AAB80263	361	LKTLDLKNNEISWTIEDMNGAFSGLDKLRRLILQGNRIRSITKKAFTGLDALEHLDLSDN
DNA37140	421	AIMSLQGNAFSQMKKLQQLHLNTSSLLCDCQLKWLPQWVAENNFQSFVNASCAHPQLLKG
P_AAB80263	421	AIMSLQGNAFSQMKKLQQLHLNTSSLLCDCQLKWLPQWVAENNFQSFVNASCAHPQLLKG
DNA37140	481	RSIFAVSPDGFVCDDFPKPQITVQPETQSAIKGSNLSFICSAASSSDSPMTFAWKKDNEL
P_AAB80263	481	RSIFAVSPDGFVCDDFPKPQITVQPETQSAIKGSNLSFICSAASSSDSPMTFAWKKDNEL
DNA37140	541	LHDAEMENYAHLRAQGGEVMEYTTILRLREVEFASEGKYQCVISNHFGSSYSVKAKLTVN ************************************
P_AAB80263	541	LHDAEMENYAHLRAQGGEVMEYTTILRLREVEFASEGKYQCVISNHFGSSYSVKAKLTVN
DNA37140	601	MLPSFTKTPMDLTIRAGAMARLECAAVGHPAPQIAWQKDGGTDFPAARERRMHVMPEDDV
P_AAB80263	601	MLPSFTKTPMDLTIRAGAMARLECAAVGHPAPQIAWQKDGGTDFPAARERRMHVMPEDDV
DNA37140	661	FFIVDVKIEDIGVYSCTAQNSAGSISANATLTVLETPSFLRPLLDRTVTKGETAVLQCIA ************************************
P_AAB80263	661	FFIVDVKIEDIGVYSCTAQNSAGSISANATLTVLETPSFLRPLLDRTVTKGETAVLQCIA
DNA37140	721	GGSPPPKLNWTKDDSPLVVTERHFFAAGNQLLIIVDSDVSDAGKYTCEMSNTLGTERGNV
P_AAB80263	721	GGSPPPKLNWTKDDSPLVVTERHFFAAGNQLLIIVDSDVSDAGKYTCEMSNTLGTERGNV
DNA37140	781	RLSVIPTPTCDSPQMTAPSLDDDGWATVGVVIIAVVCCVVGTSLVWVVIIYHTRRRNEDC
P_AAB80263	781	RLSVIPTPTCDSPQMTAPSLDDDGWATVGVVIIAVVCCVVGTSLVWVVIIYHTRRRNEDC
DNA37140	841	SITNTDETNLPADIPSYLSSQGTLADRQDGYVSSESGSHHQFVTSSGAGFFLPQHDSSGT ***********************************
P_AAB80263	841	SITNTDETNLPADIPSYLSSQGTLADRQDGYVSSESGSHHQFVTSSGAGFFLPQHDSSGT
DNA37140	901	CHIDNSSEADVEAATDLFLCPFLGSTGPMYLKGNVYGSDPFETYHTGCSPDPRTVLMDHY
P_AAB80263	901	CHIDNSSEADVEAATDLFLCPFLGSTGPMYLKGNVYGSDPFETYHTGCSPDPRTVLMDHY
DNA37140	961	EPSYIKKKECYPCSHPSEESCERSFSNISWPSHVRKLLNTSYSHNEGPGMKNLCLNKSSL
P_AAB80263	961	EPSYIKKKECYPCSHPSEESCERSFSNISWPSHVRKLLNTSYSHNEGPGMKNLCLNKSSL
DNA37140	1021	DFSANPEPASVASSNSFMGTFGKALRRPHLDAYSSFGQPSDCQPRAFYLKAHSSPDLDSG
P_AAB80263	1021	DFSANPEPASVASSNSFMGTFGKALRRPHLDAYSSFGQPSDCQPRAFYLKAHSSPDLDSG

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1081 SEEDGKERTDFQEENHICTFKQTLENYRTPNFQSYDLDT
 DNA37140
              **********
P AAB80263
          1081 SEEDGKERTDFQEENHICTFKQTLENYRTPNFQSYDLDT
>3 P AAY70674 Human PRO326 protein - Homo sapiens. (1119 aa) [1 seg]
Score = 5870 (2265 bits), Expect = 0.0
 Identities = 1119/1119 (100%), Positives = 1119/1119 (100%), at 1,1-1119,1119
 DNA37140
            1 MSAPSLRARAAGLGLLLCAVLGRAGRSDSGGRGELGOPSGVAAERPCPTTCRCLGDLLDC
P AAY70674
            1 MSAPSLRARAAGLGLLLCAVLGRAGRSDSGGRGELGQPSGVAAERPCPTTCRCLGDLLDC
 DNA37140
           61 SRKRLARLPEPLPSWVARLDLSHNRLSFIKASSMSHLOSLREVKLNNNELETIPNLGPVS
              61 SRKRLARLPEPLPSWVARLDLSHNRLSFIKASSMSHLQSLREVKLNNNELETIPNLGPVS
P AAY70674
 DNA37140
          121 ANITLLSLAGNRIVEILPEHLKEFOSLETLDLSSNNISELOTAFPALOLKYLYLNSNRVT
              ******************
P AAY70674
          121 ANITLLSLAGNRIVEILPEHLKEFQSLETLDLSSNNISELQTAFPALQLKYLYLNSNRVT
 DNA37140
          181 SMEPGYFDNLANTLLVLKLNRNRISAIPPKMFKLPOLOHLELNRNKIKNVDGLTFOGLGA
              P AAY70674
          181 SMEPGYFDNLANTLLVLKLNRNRISAIPPKMFKLPQLQHLELNRNKIKNVDGLTFQGLGA
 DNA37140
          241 LKSLKMQRNGVTKLMDGAFWGLSNMEILQLDHNNLTEITKGWLYGLLMLQELHLSQNAIN
P AAY70674
          241 LKSLKMQRNGVTKLMDGAFWGLSNMEILQLDHNNLTEITKGWLYGLLMLQELHLSQNAIN
 DNA37140
          301 RISPDAWEFCQKLSELDLTFNHLSRLDDSSFLGLSLLNTLHIGNNRVSYIADCAFRGLSS
              ***********
P AAY70674
          301 RISPDAWEFCOKLSELDLTFNHLSRLDDSSFLGLSLLNTLHIGNNRVSYIADCAFRGLSS
 DNA37140
          361 LKTLDLKNNEISWTIEDMNGAFSGLDKLRRLILQGNRIRSITKKAFTGLDALEHLDLSDN
              P AAY70674
          361 LKTLDLKNNEISWTIEDMNGAFSGLDKLRRLILQGNRIRSITKKAFTGLDALEHLDLSDN
 DNA37140
          421 AIMSLQGNAFSQMKKLQQLHLNTSSLLCDCQLKWLPQWVAENNFQSFVNASCAHPQLLKG
              *****************
P AAY70674
          421 AIMSLQGNAFSQMKKLQQLHLNTSSLLCDCQLKWLPQWVAENNFQSFVNASCAHPQLLKG
 DNA37140
          481 RSIFAVSPDGFVCDDFPKPQITVQPETQSAIKGSNLSFICSAASSSDSPMTFAWKKDNEL
P AAY70674
          481 RSIFAVSPDGFVCDDFPKPQITVQPETQSAIKGSNLSFICSAASSSDSPMTFAWKKDNEL
 DNA37140
          541 LHDAEMENYAHLRAQGGEVMEYTTILRLREVEFASEGKYQCVISNHFGSSYSVKAKLTVN
              P AAY70674
          541 LHDAEMENYAHLRAQGGEVMEYTTILRLREVEFASEGKYQCVISNHFGSSYSVKAKLTVN
 DNA37140
          601 MLPSFTKTPMDLTIRAGAMARLECAAVGHPAPQIAWQKDGGTDFPAARERRMHVMPEDDV
              P AAY70674
          601 MLPSFTKTPMDLTIRAGAMARLECAAVGHPAPQIAWQKDGGTDFPAARERRMHVMPEDDV
          661 FFIVDVKIEDIGVYSCTAQNSAGSISANATLTVLETPSFLRPLLDRTVTKGETAVLQCIA
 DNA37140
              P AAY70674
          661 FFIVDVKIEDIGVYSCTAONSAGSISANATLTVLETPSFLRPLLDRTVTKGETAVLOCIA
          721 GGSPPPKLNWTKDDSPLVVTERHFFAAGNQLLIIVDSDVSDAGKYTCEMSNTLGTERGNV
 DNA37140
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721 GGSPPPKLNWTKDDSPLVVTERHFFAAGNOLLIIVDSDVSDAGKYTCEMSNTLGTERGNV
P AAY70674
          781 RLSVIPTPTCDSPQMTAPSLDDDGWATVGVVIIAVVCCVVGTSLVWVVIIYHTRRRNEDC
 DNA37140
             **********
P AAY70674
          781 RLSVIPTPTCDSPOMTAPSLDDDGWATVGVVIIAVVCCVVGTSLVWVVIIYHTRRRNEDC
          841 SITNTDETNLPADIPSYLSSQGTLADRQDGYVSSESGSHHQFVTSSGAGFFLPQHDSSGT
 DNA37140
             841 SITNTDETNLPADIPSYLSSQGTLADRQDGYVSSESGSHHQFVTSSGAGFFLPQHDSSGT
P AAY70674
          901 CHIDNSSEADVEAATDLFLCPFLGSTGPMYLKGNVYGSDPFETYHTGCSPDPRTVLMDHY
 DNA37140
             P AAY70674
          901 CHIDNSSEADVEAATDLFLCPFLGSTGPMYLKGNVYGSDPFETYHTGCSPDPRTVLMDHY
          961 EPSYIKKKECYPCSHPSEESCERSFSNISWPSHVRKLLNTSYSHNEGPGMKNLCLNKSSL
 DNA37140
             P AAY70674
          961 EPSYIKKKECYPCSHPSEESCERSFSNISWPSHVRKLLNTSYSHNEGPGMKNLCLNKSSL
         1021 DFSANPEPASVASSNSFMGTFGKALRRPHLDAYSSFGQPSDCQPRAFYLKAHSSPDLDSG
 DNA37140
             ***********
         1021 DFSANPEPASVASSNSFMGTFGKALRRPHLDAYSSFGQPSDCQPRAFYLKAHSSPDLDSG
P AAY70674
 DNA37140
         1081 SEEDGKERTDFQEENHICTFKQTLENYRTPNFQSYDLDT
             *********
P AAY70674
         1081 SEEDGKERTDFQEENHICTFKQTLENYRTPNFQSYDLDT
>4 P AAY08114 Human PRO326 protein - Homo sapiens. (1119 aa) [1 seg]
Score = 5870 (2265 bits), Expect = 0.0
Identities = 1119/1119 (100%), Positives = 1119/1119 (100%), at 1,1-1119,1119
 DNA37140
           1 MSAPSLRARAAGLGLLLCAVLGRAGRSDSGGRGELGQPSGVAAERPCPTTCRCLGDLLDC
             P AAY08114
           1 MSAPSLRARAAGLGLLLCAVLGRAGRSDSGGRGELGQPSGVAAERPCPTTCRCLGDLLDC
 DNA37140
           61 SRKRLARLPEPLPSWVARLDLSHNRLSFIKASSMSHLQSLREVKLNNNELETIPNLGPVS
           61 SRKRLARLPEPLPSWVARLDLSHNRLSFIKASSMSHLQSLREVKLNNNELETIPNLGPVS
P AAY08114
 DNA37140
          121 ANITLLSLAGNRIVEILPEHLKEFOSLETLDLSSNNISELQTAFPALQLKYLYLNSNRVT
             121 ANITLLSLAGNRIVEILPEHLKEFQSLETLDLSSNNISELQTAFPALQLKYLYLNSNRVT
P AAY08114
 DNA37140
          181 SMEPGYFDNLANTLLVLKLNRNRISAIPPKMFKLPQLQHLELNRNKIKNVDGLTFQGLGA
             P AAY08114
          181 SMEPGYFDNLANTLLVLKLNRNRISAIPPKMFKLPQLQHLELNRNKIKNVDGLTFQGLGA
          241 LKSLKMORNGVTKLMDGAFWGLSNMEILOLDHNNLTEITKGWLYGLLMLOELHLSONAIN
 DNA37140
             P AAY08114
          241 LKSLKMQRNGVTKLMDGAFWGLSNMEILQLDHNNLTEITKGWLYGLLMLQELHLSQNAIN
 DNA37140
          301 RISPDAWEFCOKLSELDLTFNHLSRLDDSSFLGLSLLNTLHIGNNRVSYIADCAFRGLSS
P AAY08114
          301 RISPDAWEFCOKLSELDLTFNHLSRLDDSSFLGLSLLNTLHIGNNRVSYIADCAFRGLSS
 DNA37140
          361 LKTLDLKNNEISWTIEDMNGAFSGLDKLRRLILQGNRIRSITKKAFTGLDALEHLDLSDN
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361 LKTLDLKNNEISWTIEDMNGAFSGLDKLRRLILOGNRIRSITKKAFTGLDALEHLDLSDN
P AAY08114
          421 AIMSLQGNAFSQMKKLQQLHLNTSSLLCDCQLKWLPQWVAENNFQSFVNASCAHPQLLKG
 DNA37140
              P AAY08114
           421 AIMSLOGNAFSOMKKLOOLHLNTSSLLCDCOLKWLPOWVAENNFOSFVNASCAHPOLLKG
           481 RSIFAVSPDGFVCDDFPKPOITVOPETOSAIKGSNLSFICSAASSSDSPMTFAWKKDNEL
 DNA37140
           481 RSIFAVSPDGFVCDDFPKPQITVQPETQSAIKGSNLSFICSAASSSDSPMTFAWKKDNEL
P AAY08114
 DNA37140
          541 LHDAEMENYAHLRAOGGEVMEYTTILRLREVEFASEGKYOCVISNHFGSSYSVKAKLTVN
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P AAY08114
          541 LHDAEMENYAHLRAQGGEVMEYTTILRLREVEFASEGKYQCVISNHFGSSYSVKAKLTVN
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          601 MLPSFTKTPMDLTIRAGAMARLECAAVGHPAPQIAWQKDGGTDFPAARERRMHVMPEDDV
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P AAY08114
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 DNA37140
          661 FFIVDVKIEDIGVYSCTAONSAGSISANATLTVLETPSFLRPLLDRTVTKGETAVLOCIA
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          661 FFIVDVKIEDIGVYSCTAONSAGSISANATLTVLETPSFLRPLLDRTVTKGETAVLQCIA
P AAY08114
 DNA37140
          721 GGSPPPKLNWTKDDSPLVVTERHFFAAGNOLLIIVDSDVSDAGKYTCEMSNTLGTERGNV
              721 GGSPPPKLNWTKDDSPLVVTERHFFAAGNOLLIIVDSDVSDAGKYTCEMSNTLGTERGNV
P AAY08114
 DNA37140
          781 RLSVIPTPTCDSPOMTAPSLDDDGWATVGVVIIAVVCCVVGTSLVWVVIIYHTRRRNEDC
              P AAY08114
          781 RLSVIPTPTCDSPQMTAPSLDDDGWATVGVVIIAVVCCVVGTSLVWVVIIYHTRRRNEDC
 DNA37140
          841 SITNTDETNLPADIPSYLSSQGTLADRQDGYVSSESGSHHQFVTSSGAGFFLPQHDSSGT
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P AAY08114
          841 SITNTDETNLPADIPSYLSSQGTLADRQDGYVSSESGSHHQFVTSSGAGFFLPQHDSSGT
 DNA37140
          901 CHIDNSSEADVEAATDLFLCPFLGSTGPMYLKGNVYGSDPFETYHTGCSPDPRTVLMDHY
              ************
P AAY08114
          901 CHIDNSSEADVEAATDLFLCPFLGSTGPMYLKGNVYGSDPFETYHTGCSPDPRTVLMDHY
 DNA37140
          961 EPSYTKKKECYPCSHPSEESCERSFSNISWPSHVRKLLNTSYSHNEGPGMKNLCLNKSSL
              P AAY08114
          961 EPSYIKKKECYPCSHPSEESCERSFSNISWPSHVRKLLNTSYSHNEGPGMKNLCLNKSSL
         1021 DFSANPEPASVASSNSFMGTFGKALRRPHLDAYSSFGOPSDCOPRAFYLKAHSSPDLDSG
 DNA37140
              P AAY08114
         1021 DFSANPEPASVASSNSFMGTFGKALRRPHLDAYSSFGQPSDCQPRAFYLKAHSSPDLDSG
 DNA37140
         1081 SEEDGKERTDFQEENHICTFKQTLENYRTPNFQSYDLDT
P AAY08114
         1081 SEEDGKERTDFQEENHICTFKQTLENYRTPNFQSYDLDT
>5 P AAY13395 protein PRO326 - Homo sapiens. (1120 aa) [1 seg]
Score = 5858 (2261 bits), Expect = 0.0
Identities = 1119/1120 (99\%), Positives = 1119/1120 (99\%), Gaps = 1/1120 (0\%),
at 1,1-1119,1120
 DNA37140
            1 MSAPSLRARAAGLGLLLCAVLGRAGRSDSGGRGELGQPSGVAAERPCPTTCRCLGDLLDC
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P_AAY13395	. 1	${\tt MSAPSLRARAAGLGLLLCAVLGRAGRSDSGGRGELGQPSGVAAERPCPTTCRCLGDLLDC}$
DNA37140	61	SRKRLARLPEPLPSWVARLDLSHNRLSFIKASSMSHLQSLREVKLNNNELETIPNLGPVS ************************************
P_AAY13395	61	SRKRLARLPEPLPSWVARLDLSHNRLSFIKASSMSHLQSLREVKLNNNELETIPNLGPVS
DNA37140	121	ANITLLSLAGNRIVEILPEHLKEFQSLETLDLSSNNISELQTAFPALQLKYLYLNSNRVT
P_AAY13395	121	ANITLLSLAGNRIVEILPEHLKEFQSLETLDLSSNNISELQTAFPALQLKYLYLNSNRVT
DNA37140	181	SMEPGYFDNLANTLLVLKLNRNRISAIPPKMFKLPQLQHLELNRNKIKNVDGLTFQGLGA
P_AAY13395	181	SMEPGYFDNLANTLLVLKLNRNRISAIPPKMFKLPQLQHLELNRNKIKNVDGLTFQGLGA
DNA37140	241	LKSLKMQRNGVTKLMDGAFWGLSNMEILQLDHNNLTEITKGWLYGLLMLQELHLSQNAIN ***********************************
P_AAY13395	241	LKSLKMQRNGVTKLMDGAFWGLSNMEILQLDHNNLTEITKGWLYGLLMLQELHLSQNAIN
DNA37140	301	RISPDAWEFCQKLSELDLTFNHLSRLDDSSFLGLSLLNTLHIGNNRVSYIADCAFRGLSS ***********************************
P_AAY13395	301	RISPDAWEFCQKLSELDLTFNHLSRLDDSSFLGLSLLNTLHIGNNRVSYIADCAFRGLSS
DNA37140	361	LKTLDLKNNEISWTIEDMNGAFSGLDKLRRLILQGNRIRSITKKAFTGLDALEHLDLSDN ************************************
P_AAY13395	361	LKTLDLKNNEISWTIEDMNGAFSGLDKLRRLILQGNRIRSITKKAFTGLDALEHLDLSDN
DNA37140	421	AIMSLQGNAFSQMKKLQQLHLNTSSLLCDCQLKWLPQWVAENNFQSFVNASCAHPQLLKG
P_AAY13395	421	AIMSLQGNAFSQMKKLQQLHLNTSSLLCDCQLKWLPQWVAENNFQSFVNASCAHPQLLKG
DNA37140	481	RSIFAVSPDGFVCDDFPKPQITVQPETQSAIKGSNLSFICSAASSSDSPMTFAWKKDNEL ************************************
P_AAY13395	481	RSIFAVSPDGFVCDDFPKPQITVQPETQSAIKGSNLSFICSAASSSDSPMTFAWKKDNEL
DNA37140	541	LHDAEMENYAHLRAQGGEVMEYTTILRLREVEFASEGKYQCVISNHFGSSYSVKAKLTVN ************************************
P_AAY13395	541	LHDAEMENYAHLRAQGGEVMEYTTILRLREVEFASEGKYQCVISNHFGSSYSVKAKLTVN
DNA37140	601	MLPSFTKTPMDLTIRAGAMARLECAAVGHPAPQIAWQKDGGTDFPAARERRMHVMPEDDV ***********************************
P_AAY13395	601	MLPSFTKTPMDLTIRAGAMARLECAAVGHPAPQIAWQKDGGTDFPAARERRMHVMPEDDV
DNA37140	661	FFIVDVKIEDIGVYSCTAQNSAGSISANATLTVLETPSFLRPLLDRTVTKGETAVLQCIA
P_AAY13395	661	FFIVDVKIEDIGVYSCTAQNSAGSISANATLTVLETPSFLRPLLDRTVTKGETAVLQCIA
DNA37140	721	-GGSPPPKLNWTKDDSPLVVTERHFFAAGNQLLIIVDSDVSDAGKYTCEMSNTLGTERGN
P_AAY13395	721	QGGSPPPKLNWTKDDSPLVVTERHFFAAGNQLLIIVDSDVSDAGKYTCEMSNTLGTERGN
DNA37140	780	VRLSVIPTPTCDSPQMTAPSLDDDGWATVGVVIIAVVCCVVGTSLVWVVIIYHTRRRNED
P_AAY13395	781	VRLSVIPTPTCDSPQMTAPSLDDDGWATVGVVIIAVVCCVVGTSLVWVVIIYHTRRRNED
DNA37140	840	CSITNTDETNLPADIPSYLSSQGTLADRQDGYVSSESGSHHQFVTSSGAGFFLPQHDSSG
P_AAY13395	841	CSITNTDETNLPADIPSYLSSQGTLADRQDGYVSSESGSHHQFVTSSGAGFFLPQHDSSG

DNA37140	900	TCHIDNSSEADVEAATDLFLCPFLGSTGPMYLKGNVYGSDPFETYHTGCSPDPRTVLMDH
P_AAY13395	901	TCHIDNSSEADVEAATDLFLCPFLGSTGPMYLKGNVYGSDPFETYHTGCSPDPRTVLMDH
DNA37140	960	YEPSYIKKKECYPCSHPSEESCERSFSNISWPSHVRKLLNTSYSHNEGPGMKNLCLNKSS ***********************************
P_AAY13395	961	YEPSYIKKKECYPCSHPSEESCERSFSNISWPSHVRKLLNTSYSHNEGPGMKNLCLNKSS
DNA37140	1020	LDFSANPEPASVASSNSFMGTFGKALRRPHLDAYSSFGQPSDCQPRAFYLKAHSSPDLDS ***********************************
P_AAY13395	1021	LDFSANPEPASVASSNSFMGTFGKALRRPHLDAYSSFGQPSDCQPRAFYLKAHSSPDLDS
DNA37140	1080	GSEEDGKERTDFQEENHICTFKQTLENYRTPNFQSYDLDT ***********************************
P_AAY13395	1081	GSEEDGKERTDFQEENHICTFKQTLENYRTPNFQSYDLDT
Score = 54	62 (23	man PRO335 protein - Homo sapiens. (1059 aa) [1 seg] 108 bits), Expect = 0.0 41/1044 (99%), Positives = 1043/1044 (99%), at 76,16-1119,1059
DNA37140		VARLDLSHNRLSFIKASSMSHLQSLREVKLNNNELETIPNLGPVSANITLLSLAGNRIVE
		* *************
P_AAB80261		ISRPDLSHNRLSFIKASSMSHLQSLREVKLNNNELETIPNLGPVSANITLLSLAGNRIVE
DNA37140	136	ILPEHLKEFQSLETLDLSSNNISELQTAFPALQLKYLYLNSNRVTSMEPGYFDNLANTLL ***********************************
P_AAB80261	76	ILPEHLKEFQSLETLDLSSNNISELQTAFPALQLKYLYLNSNRVTSMEPGYFDNLANTLL
DNA37140	196	VLKLNRNRISAIPPKMFKLPQLQHLELNRNKIKNVDGLTFQGLGALKSLKMQRNGVTKLM ************************************
P_AAB80261	136	VLKLNRNRISAIPPKMFKLPQLQHLELNRNKIKNVDGLTFQGLGALKSLKMQRNGVTKLM
DNA37140	256	DGAFWGLSNMEILQLDHNNLTEITKGWLYGLLMLQELHLSQNAINRISPDAWEFCQKLSE
P_AAB80261	196	DGAFWGLSNMEILQLDHNNLTEITKGWLYGLLMLQELHLSQNAINRISPDAWEFCQKLSE
DNA37140	316	LDLTFNHLSRLDDSSFLGLSLLNTLHIGNNRVSYIADCAFRGLSSLKTLDLKNNEISWTI
P_AAB80261	256	LDLTFNHLSRLDDSSFLGLSLLNTLHIGNNRVSYIADCAFRGLSSLKTLDLKNNEISWTI
DNA37140	376	EDMNGAFSGLDKLRRLILQGNRIRSITKKAFTGLDALEHLDLSDNAIMSLQGNAFSQMKK
P_AAB80261	316	EDMNGAFSGLDKLRRLILQGNRIRSITKKAFTGLDALEHLDLSDNAIMSLQGNAFSQMKK
DNA37140	436	LQQLHLNTSSLLCDCQLKWLPQWVAENNFQSFVNASCAHPQLLKGRSIFAVSPDGFVCDD
P_AAB80261	376	LQQLHLNTSSLLCDCQLKWLPQWVAENNFQSFVNASCAHPQLLKGRSIFAVSPDGFVCDD
DNA37140	496	FPKPQITVQPETQSAIKGSNLSFICSAASSSDSPMTFAWKKDNELLHDAEMENYAHLRAQ
P_AAB80261	436	FPKPQITVQPETQSAIKGSNLSFICSAASSSDSPMTFAWKKDNELLHDAEMENYAHLRAQ
DNA37140	556	GGEVMEYTTILRLREVEFASEGKYQCVISNHFGSSYSVKAKLTVNMLPSFTKTPMDLTIR
P_AAB80261	496	GGEVMEYTTILRLREVEFASEGKYQCVISNHFGSSYSVKAKLTVNMLPSFTKTPMDLTIR

DNA37140	616	AGAMARLECAAVGHPAPQIAWQKDGGTDFPAARERRMHVMPEDDVFFIVDVKIEDIGVYS
P_AAB80261	556	AGAMARLECAAVGHPAPQIAWQKDGGTDFPAARERRMHVMPEDDVFFIVDVKIEDIGVYS
DNA37140	676	CTAQNSAGSISANATLTVLETPSFLRPLLDRTVTKGETAVLQCIAGGSPPPKLNWTKDDS
P_AAB80261	616	CTAQNSAGSISANATLTVLETPSFLRPLLDRTVTKGETAVLQCIAGGSPPPKLNWTKDDS
DNA37140	736	PLVVTERHFFAAGNQLLIIVDSDVSDAGKYTCEMSNTLGTERGNVRLSVIPTPTCDSPQM
P_AAB80261	676	PLVVTERHFFAAGNQLLIIVDSDVSDAGKYTCEMSNTLGTERGNVRLSVIPTPTCDSPQM
DNA37140	796	TAPSLDDDGWATVGVVIIAVVCCVVGTSLVWVVIIYHTRRRNEDCSITNTDETNLPADIP
P_AAB80261	736	TAPSLDDDGWATVGVVIIAVVCCVVGTSLVWVVIIYHTRRRNEDCSITNTDETNLPADIP
DNA37140	856	SYLSSQGTLADRQDGYVSSESGSHHQFVTSSGAGFFLPQHDSSGTCHIDNSSEADVEAAT
P_AAB80261	796	SYLSSQGTLADRQDGYVSSESGSHHQFVTSSGAGFFLPQHDSSGTCHIDNSSEADVEAAT
DNA37140	916	DLFLCPFLGSTGPMYLKGNVYGSDPFETYHTGCSPDPRTVLMDHYEPSYIKKKECYPCSH
P_AAB80261	856	DLFLCPFLGSTGPMYLKGNVYGSDPFETYHTGCSPDPRTVLMDHYEPSYIKKKECYPCSH
DNA37140	976	PSEESCERSFSNISWPSHVRKLLNTSYSHNEGPGMKNLCLNKSSLDFSANPEPASVASSN ***********************************
P_AAB80261	916	PSEESCERSFSNISWPSHVRKLLNTSYSHNEGPGMKNLCLNKSSLDFSANPEPASVASSN
DNA37140	1036	SFMGTFGKALRRPHLDAYSSFGQPSDCQPRAFYLKAHSSPDLDSGSEEDGKERTDFQEEN
P_AAB80261	976	SFMGTFGKALRRPHLDAYSSFGQPSDCQPRAFYLKAHSSPDLDSGSEEDGKERTDFQEEN
DNA37140	1096	HICTFKQTLENYRTPNFQSYDLDT ***********************************
P_AAB80261	1036	HICTFKQTLENYRTPNFQSYDLDT
		nan PRO335 protein - Homo sapiens. (1059 aa) [1 seg] 108 bits), Expect = 0.0
		11/1044 (99%), Positives = 1043/1044 (99%), at 76,16-1119,1059
DNA37140	76	VARLDLSHNRLSFIKASSMSHLQSLREVKLNNNELETIPNLGPVSANITLLSLAGNRIVE* **********************************
P_AAY70672	16	ISRPDLSHNRLSFIKASSMSHLQSLREVKLNNNELETIPNLGPVSANITLLSLAGNRIVE
DNA37140	136	ILPEHLKEFQSLETLDLSSNNISELQTAFPALQLKYLYLNSNRVTSMEPGYFDNLANTLL
P_AAY70672	76	ILPEHLKEFQSLETLDLSSNNISELQTAFPALQLKYLYLNSNRVTSMEPGYFDNLANTLL
DNA37140	196	VLKLNRNRISAIPPKMFKLPQLQHLELNRNKIKNVDGLTFQGLGALKSLKMQRNGVTKLM ************************************
P_AAY70672	136	VLKLNRNRISAIPPKMFKLPQLQHLELNRNKIKNVDGLTFQGLGALKSLKMQRNGVTKLM
DNA37140	256	DGAFWGLSNMEILQLDHNNLTEITKGWLYGLLMLQELHLSQNAINRISPDAWEFCQKLSE
P_AAY70672	196	DGAFWGLSNMEILQLDHNNLTEITKGWLYGLLMLQELHLSQNAINRISPDAWEFCQKLSE
DNA37140	316	LDLTFNHLSRLDDSSFLGLSLLNTLHIGNNRVSYIADCAFRGLSSLKTLDLKNNEISWTI

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***********
          256 LDLTFNHLSRLDDSSFLGLSLLNTLHIGNNRVSYIADCAFRGLSSLKTLDLKNNEISWTI
P AAY70672
          376 EDMNGAFSGLDKLRRLILOGNRIRSITKKAFTGLDALEHLDLSDNAIMSLQGNAFSQMKK
 DNA37140
             ***********
          316 EDMNGAFSGLDKLRRLILQGNRIRSITKKAFTGLDALEHLDLSDNAIMSLQGNAFSQMKK
P AAY70672
          436 LOOLHLNTSSLLCDCQLKWLPQWVAENNFQSFVNASCAHPQLLKGRSIFAVSPDGFVCDD
 DNA37140
              **************
          376 LOOLHLNTSSLLCDCQLKWLPQWVAENNFQSFVNASCAHPQLLKGRSIFAVSPDGFVCDD
P AAY70672
          496 FPKPOITVOPETQSAIKGSNLSFICSAASSSDSPMTFAWKKDNELLHDAEMENYAHLRAQ
 DNA37140
                    436 FPKPQITVQPETQSAIKGSNLSFICSAASSSDSPMTFAWKKDNELLHDAEMENYAHLRAQ
P AAY70672
          556 GGEVMEYTTILRLREVEFASEGKYQCVISNHFGSSYSVKAKLTVNMLPSFTKTPMDLTIR
 DNA37140
             P AAY70672
          496 GGEVMEYTTILRLREVEFASEGKYOCVISNHFGSSYSVKAKLTVNMLPSFTKTPMDLTIR
          616 AGAMARLECAAVGHPAPQIAWQKDGGTDFPAARERRMHVMPEDDVFFIVDVKIEDIGVYS
 DNA37140
             556 AGAMARLECAAVGHPAPOIAWOKDGGTDFPAARERRMHVMPEDDVFFIVDVKIEDIGVYS
P AAY70672
          676 CTAONSAGSISANATLTVLETPSFLRPLLDRTVTKGETAVLQCIAGGSPPPKLNWTKDDS
 DNA37140
             ***********
          616 CTAONSAGSISANATLTVLETPSFLRPLLDRTVTKGETAVLQCIAGGSPPPKLNWTKDDS
P AAY70672
          736 PLVVTERHFFAAGNQLLIIVDSDVSDAGKYTCEMSNTLGTERGNVRLSVIPTPTCDSPQM
 DNA37140
             676 PLVVTERHFFAAGNQLLIIVDSDVSDAGKYTCEMSNTLGTERGNVRLSVIPTPTCDSPQM
P AAY70672
          796 TAPSLDDDGWATVGVVIIAVVCCVVGTSLVWVVIIYHTRRRNEDCSITNTDETNLPADIP
 DNA37140
             736 TAPSLDDDGWATVGVVIIAVVCCVVGTSLVWVVIIYHTRRRNEDCSITNTDETNLPADIP
P AAY70672
          856 SYLSSQGTLADRQDGYVSSESGSHHQFVTSSGAGFFLPQHDSSGTCHIDNSSEADVEAAT
 DNA37140
             ***********
          796 SYLSSQGTLADRQDGYVSSESGSHHQFVTSSGAGFFLPQHDSSGTCHIDNSSEADVEAAT
P AAY70672
          916 DLFLCPFLGSTGPMYLKGNVYGSDPFETYHTGCSPDPRTVLMDHYEPSYIKKKECYPCSH
 DNA37140
          856 DLFLCPFLGSTGPMYLKGNVYGSDPFETYHTGCSPDPRTVLMDHYEPSYIKKKECYPCSH
P AAY70672
          976 PSEESCERSFSNISWPSHVRKLLNTSYSHNEGPGMKNLCLNKSSLDFSANPEPASVASSN
 DNA37140
             ************
          916 PSEESCERSFSNISWPSHVRKLLNTSYSHNEGPGMKNLCLNKSSLDFSANPEPASVASSN
P AAY70672
         1036 SFMGTFGKALRRPHLDAYSSFGQPSDCQPRAFYLKAHSSPDLDSGSEEDGKERTDFQEEN
 DNA37140
             976 SFMGTFGKALRRPHLDAYSSFGQPSDCQPRAFYLKAHSSPDLDSGSEEDGKERTDFQEEN
P AAY70672
         1096 HICTFKOTLENYRTPNFQSYDLDT
 DNA37140
P AAY70672
        1036 HICTFKQTLENYRTPNFQSYDLDT
>8 P AAY13393 protein PRO335 - Homo sapiens. (1059 aa) [1 seg]
Score = 5462 (2108 bits), Expect = 0.0
```

Identities	1041/1044 (99%), Positives = 1043/1044 (99%), at 76,16-1119,1059
DNA37140	76 VARLDLSHNRLSFIKASSMSHLQSLREVKLNNNELETIPNLGPVSANITLLSLAGNRIVE
P_AAY13393	16 ISRPDLSHNRLSFIKASSMSHLQSLREVKLNNNELETIPNLGPVSANITLLSLAGNRIVE
DNA37140	136 ILPEHLKEFQSLETLDLSSNNISELQTAFPALQLKYLYLNSNRVTSMEPGYFDNLANTLL
P_AAY13393	76 ILPEHLKEFQSLETLDLSSNNISELQTAFPALQLKYLYLNSNRVTSMEPGYFDNLANTLL
DNA37140	196 VLKLNRNRISAIPPKMFKLPQLQHLELNRNKIKNVDGLTFQGLGALKSLKMQRNGVTKLM
P_AAY13393	136 VLKLNRNRISAIPPKMFKLPQLQHLELNRNKIKNVDGLTFQGLGALKSLKMQRNGVTKLM
DNA37140	256 DGAFWGLSNMEILQLDHNNLTEITKGWLYGLLMLQELHLSQNAINRISPDAWEFCQKLSE
P_AAY13393	196 DGAFWGLSNMEILQLDHNNLTEITKGWLYGLLMLQELHLSQNAINRISPDAWEFCQKLSE
DNA37140	316 LDLTFNHLSRLDDSSFLGLSLLNTLHIGNNRVSYIADCAFRGLSSLKTLDLKNNEISWTI
P_AAY13393	256 LDLTFNHLSRLDDSSFLGLSLLNTLHIGNNRVSYIADCAFRGLSSLKTLDLKNNEISWTI
DNA37140	376 EDMNGAFSGLDKLRRLILQGNRIRSITKKAFTGLDALEHLDLSDNAIMSLQGNAFSQMKK
P_AAY13393	316 EDMNGAFSGLDKLRRLILQGNRIRSITKKAFTGLDALEHLDLSDNAIMSLQGNAFSQMKK
DNA37140	436 LQQLHLNTSSLLCDCQLKWLPQWVAENNFQSFVNASCAHPQLLKGRSIFAVSPDGFVCDD
P_AAY13393	376 LQQLHLNTSSLLCDCQLKWLPQWVAENNFQSFVNASCAHPQLLKGRSIFAVSPDGFVCDD
DNA37140	496 FPKPQITVQPETQSAIKGSNLSFICSAASSSDSPMTFAWKKDNELLHDAEMENYAHLRAQ
P_AAY13393	436 FPKPQITVQPETQSAIKGSNLSFICSAASSSDSPMTFAWKKDNELLHDAEMENYAHLRAQ
DNA37140	556 GGEVMEYTTILRLREVEFASEGKYQCVISNHFGSSYSVKAKLTVNMLPSFTKTPMDLTIR
P_AAY13393	496 GGEVMEYTTILRLREVEFASEGKYQCVISNHFGSSYSVKAKLTVNMLPSFTKTPMDLTIR
DNA37140	616 AGAMARLECAAVGHPAPQIAWQKDGGTDFPAARERRMHVMPEDDVFFIVDVKIEDIGVYS
P_AAY13393	556 AGAMARLECAAVGHPAPQIAWQKDGGTDFPAARERRMHVMPEDDVFFIVDVKIEDIGVYS
DNA37140	676 CTAQNSAGSISANATLTVLETPSFLRPLLDRTVTKGETAVLQCIAGGSPPPKLNWTKDDS
P_AAY13393	616 CTAQNSAGSISANATLTVLETPSFLRPLLDRTVTKGETAVLQCIAGGSPPPKLNWTKDDS
DNA37140	736 PLVVTERHFFAAGNQLLIIVDSDVSDAGKYTCEMSNTLGTERGNVRLSVIPTPTCDSPQM ************************************
P_AAY13393	676 PLVVTERHFFAAGNQLLIIVDSDVSDAGKYTCEMSNTLGTERGNVRLSVIPTPTCDSPQM
DNA37140	796 TAPSLDDDGWATVGVVIIAVVCCVVGTSLVWVVIIYHTRRRNEDCSITNTDETNLPADIP ************************************
P_AAY13393	736 TAPSLDDDGWATVGVVIIAVVCCVVGTSLVWVVIIYHTRRRNEDCSITNTDETNLPADIP
DNA37140	856 SYLSSQGTLADRQDGYVSSESGSHHQFVTSSGAGFFLPQHDSSGTCHIDNSSEADVEAAT ***********************************
P_AAY13393	796 SYLSSQGTLADRQDGYVSSESGSHHQFVTSSGAGFFLPQHDSSGTCHIDNSSEADVEAAT

DNA37140	916	DLFLCPFLGSTGPMYLKGNVYGSDPFETYHTGCSPDPRTVLMDHYEPSYIKKKECYPCSH ************************************
P_AAY13393	856	DLFLCPFLGSTGPMYLKGNVYGSDPFETYHTGCSPDPRTVLMDHYEPSYIKKKECYPCSH
DNA37140	976	PSEESCERSFSNISWPSHVRKLLNTSYSHNEGPGMKNLCLNKSSLDFSANPEPASVASSN
P_AAY13393	916	PSEESCERSFSNISWPSHVRKLLNTSYSHNEGPGMKNLCLNKSSLDFSANPEPASVASSN
DNA37140	1036	SFMGTFGKALRRPHLDAYSSFGQPSDCQPRAFYLKAHSSPDLDSGSEEDGKERTDFQEEN
P_AAY13393	976	SFMGTFGKALRRPHLDAYSSFGQPSDCQPRAFYLKAHSSPDLDSGSEEDGKERTDFQEEN
DNA37140	1096	HICTFKQTLENYRTPNFQSYDLDT
P_AAY13393	1036	HICTFKQTLENYRTPNFQSYDLDT
Score = 54	62 (2)	man PRO335 protein - Homo sapiens. (1059 aa) [1 seg] 108 bits), Expect = 0.0 41/1044 (99%), Positives = 1043/1044 (99%), at 76,16-1119,1059
DNA37140		
		VARLDLSHNRLSFIKASSMSHLQSLREVKLNNNELETIPNLGPVSANITLLSLAGNRIVE* **********************************
P_AAY08095	16	ISRPDLSHNRLSFIKASSMSHLQSLREVKLNNNELETIPNLGPVSANITLLSLAGNRIVE
DNA37140	136	ILPEHLKEFQSLETLDLSSNNISELQTAFPALQLKYLYLNSNRVTSMEPGYFDNLANTLL ***********************************
P_AAY08095	76	ILPEHLKEFQSLETLDLSSNNISELQTAFPALQLKYLYLNSNRVTSMEPGYFDNLANTLL
DNA37140	196	VLKLNRNRISAIPPKMFKLPQLQHLELNRNKIKNVDGLTFQGLGALKSLKMQRNGVTKLM
P_AAY08095	136	VLKLNRNRISAIPPKMFKLPQLQHLELNRNKIKNVDGLTFQGLGALKSLKMQRNGVTKLM
DNA37140	256	DGAFWGLSNMEILQLDHNNLTEITKGWLYGLLMLQELHLSQNAINRISPDAWEFCQKLSE ************************************
P_AAY08095	196	DGAFWGLSNMEILQLDHNNLTEITKGWLYGLLMLQELHLSQNAINRISPDAWEFCQKLSE
DNA37140	316	LDLTFNHLSRLDDSSFLGLSLLNTLHIGNNRVSYIADCAFRGLSSLKTLDLKNNEISWTI
P_AAY08095	256	LDLTFNHLSRLDDSSFLGLSLLNTLHIGNNRVSYIADCAFRGLSSLKTLDLKNNEISWTI
DNA37140	376	EDMNGAFSGLDKLRRLILQGNRIRSITKKAFTGLDALEHLDLSDNAIMSLQGNAFSQMKK
P_AAY08095	316	EDMNGAFSGLDKLRRLILQGNRIRSITKKAFTGLDALEHLDLSDNAIMSLQGNAFSQMKK
DNA37140	436	LQQLHLNTSSLLCDCQLKWLPQWVAENNFQSFVNASCAHPQLLKGRSIFAVSPDGFVCDD
P_AAY08095	376	LQQLHLNTSSLLCDCQLKWLPQWVAENNFQSFVNASCAHPQLLKGRSIFAVSPDGFVCDD
DNA37140	496	FPKPQITVQPETQSAIKGSNLSFICSAASSSDSPMTFAWKKDNELLHDAEMENYAHLRAQ
P_AAY08095	436	FPKPQITVQPETQSAIKGSNLSFICSAASSSDSPMTFAWKKDNELLHDAEMENYAHLRAQ
DNA37140	556	GGEVMEYTTILRLREVEFASEGKYQCVISNHFGSSYSVKAKLTVNMLPSFTKTPMDLTIR
P_AAY08095	496	**************************************

DNA37140	616	AGAMARLECAAVGHPAPQIAWQKDGGTDFPAARERRMHVMPEDDVFFIVDVKIEDIGVYS ************************************
P_AAY08095	556	AGAMARLECAAVGHPAPQIAWQKDGGTDFPAARERRMHVMPEDDVFFIVDVKIEDIGVYS
DNA37140	676	CTAQNSAGSISANATLTVLETPSFLRPLLDRTVTKGETAVLQCIAGGSPPPKLNWTKDDS ***********************************
P_AAY08095	616	CTAQNSAGSISANATLTVLETPSFLRPLLDRTVTKGETAVLQCIAGGSPPPKLNWTKDDS
DNA37140	736	PLVVTERHFFAAGNQLLIIVDSDVSDAGKYTCEMSNTLGTERGNVRLSVIPTPTCDSPQM ************************************
P_AAY08095	676	PLVVTERHFFAAGNQLLIIVDSDVSDAGKYTCEMSNTLGTERGNVRLSVIPTPTCDSPQM
DNA37140	796	TAPSLDDDGWATVGVVIIAVVCCVVGTSLVWVVIIYHTRRRNEDCSITNTDETNLPADIP ************************************
P_AAY08095	736	TAPSLDDDGWATVGVVIIAVVCCVVGTSLVWVVIIYHTRRRNEDCSITNTDETNLPADIP
DNA37140	856	SYLSSQGTLADRQDGYVSSESGSHHQFVTSSGAGFFLPQHDSSGTCHIDNSSEADVEAAT ***********************************
P_AAY08095	796	SYLSSQGTLADRQDGYVSSESGSHHQFVTSSGAGFFLPQHDSSGTCHIDNSSEADVEAAT
DNA37140	916	DLFLCPFLGSTGPMYLKGNVYGSDPFETYHTGCSPDPRTVLMDHYEPSYIKKKECYPCSH ************************************
P_AAY08095	856	DLFLCPFLGSTGPMYLKGNVYGSDPFETYHTGCSPDPRTVLMDHYEPSYIKKKECYPCSH
DNA37140	976	PSEESCERSFSNISWPSHVRKLLNTSYSHNEGPGMKNLCLNKSSLDFSANPEPASVASSN ***********************************
P_AAY08095	916	PSEESCERSFSNISWPSHVRKLLNTSYSHNEGPGMKNLCLNKSSLDFSANPEPASVASSN
DNA37140	1036	SFMGTFGKALRRPHLDAYSSFGQPSDCQPRAFYLKAHSSPDLDSGSEEDGKERTDFQEEN ***********************************
P_AAY08095	976	SFMGTFGKALRRPHLDAYSSFGQPSDCQPRAFYLKAHSSPDLDSGSEEDGKERTDFQEEN
DNA37140	1096	HICTFKQTLENYRTPNFQSYDLDT ***********************************
P_AAY08095	1036	HICTFKQTLENYRTPNFQSYDLDT

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3481 TCTTACGTAAATTAATGATATAAATCATGATTATTTTATGTATTTTTATAATGCCAGATT
P AAF72422
         3903 TCTTTTTATGGAAAATGAGTTACTAAAGCATTTTAAATAATACCTGCCTTGTACCATTTT
 DNA37140
             *****************
         3541 TCTTTTTATGGAAAATGAGTTACTAAAGCATTTTAAATAATACCTGCCTTGTACCATTTT
P AAF72422
         3963 TTAAATAGAAGTTACTTCATTATATTTTGCACATTATATTTAATAAAATGTGTCAATTTG
 DNA37140
         3601 TTAAATAGAAGTTACTTCATTATATTTTGCACATTATATTTAATAAAATGTGTCAATTTG
P AAF72422
 DNA37140
         4023 AA
P AAF72422
         3661 AA
>9 P AAZ52206 Human PRO335 protein encoding cDNA, UNQ287V. DNA, PAT 18-JUL-2000
(3662 bp) [1 seg]
Score = 3542 (7022 bits), Expect = 0.0
Identities = 3542/3542 (100%), at 483,121-4024,3662, Strand +/+
          483 GGACTTAAGTCACAACAGATTATCTTTCATCAAGGCAAGTTCCATGAGCCACCTTCAAAG
 DNA37140
             ******************
          121 GGACTTAAGTCACAACAGATTATCTTTCATCAAGGCAAGTTCCATGAGCCACCTTCAAAG
P AAZ52206
          543 CCTTCGAGAAGTGAAACTGAACAATGAATTGGAGACCATTCCAAATCTGGGACCAGT
 DNA37140
             ******************
          181 CCTTCGAGAAGTGAAACTGAACAATGAATTGGAGACCATTCCAAATCTGGGACCAGT
P AAZ52206
          603 CTCGGCAAATATTACACTTCTCCTTGGCTGGAAACAGGATTGTTGAAATACTCCCTGA
 DNA37140
             ******************
          241 CTCGGCAAATATTACACTTCTCCTTGGCTGGAAACAGGATTGTTGAAATACTCCCTGA
P AAZ52206
          663 ACATCTGAAAGAGTTTCAGTCCCTTGAAACTTTGGACCTTAGCAGCAACAATATTTCAGA
 DNA37140
             *************
          301 ACATCTGAAAGAGTTTCAGTCCCTTGAAACTTTGGACCTTAGCAGCAACAATATTTCAGA
P AAZ52206
          723 GCTCCAAACTGCATTTCCAGCCCTACAGCTCAAATATCTGTATCTCAACAGCAACCGAGT
 DNA37140
             ****************
          361 GCTCCAAACTGCATTTCCAGCCCTACAGCTCAAATATCTGTATCTCAACAGCAACCGAGT
P AAZ52206
          783 CACATCAATGGAACCTGGGTATTTTGACAATTTGGCCAACACACTCCTTGTGTTAAAGCT
 DNA37140
             421 CACATCAATGGAACCTGGGTATTTTGACAATTTGGCCAACACACTCCTTGTGTTAAAGCT
P AAZ52206
          843 GAACAGGAACCGAATCTCAGCTATCCCACCCAAGATGTTTAAACTGCCCCAACTGCAACA
 DNA37140
              ************
          481 GAACAGGAACCGAATCTCAGCTATCCCACCCAAGATGTTTAAACTGCCCCAACTGCAACA
P AAZ52206
          903 TCTCGAATTGAACCGAAACAAGATTAAAAATGTAGATGGACTGACATTCCAAGGCCTTGG
 DNA37140
             *****************
          541 TCTCGAATTGAACCGAAACAAGATTAAAAATGTAGATGGACTGACATTCCAAGGCCTTGG
P AAZ52206
          DNA37140
             P AAZ52206
         1023 TTGGGGGCTGAGCAACATGGAAATTTTGCAGCTGGACCATAACAACCTAACAGAGATTAC
 DNA37140
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P_AAZ52206	661	TTGGGGGCTGAGCAACATGGAAATTTTGCAGCTGGACCATAACAACCTAACAGAGATTAC	
DNA37140	1083	CAAAGGCTGGCTTTACGGCTTGCTGATGCTGCAGGAACTTCATCTCAGCCAAAATGCCAT	
P_AAZ52206	721	CAAAGGCTGGCTTTACGGCTTGCTGATGCTGCAGGAACTTCATCTCAGCCAAAATGCCAT	
DNA37140	1143	CAACAGGATCAGCCCTGATGCCTGGGAGTTCTGCCAGAAGCTCAGTGAGCTGGACCTAAC	
P_AAZ52206	781	CAACAGGATCAGCCCTGATGCCTGGGAGTTCTGCCAGAAGCTCAGTGAGCTGGACCTAAC	
DNA37140	1203	TTTCAATCACTTATCAAGGTTAGATGATTCAAGCTTCCTTGGCCTAAGCTTACTAAATAC	
P_AAZ52206	841	TTTCAATCACTTATCAAGGTTAGATGATTCAAGCTTCCTTGGCCTAAGCTTACTAAATAC	
DNA37140	1263	ACTGCACATTGGGAACAACAGAGTCAGCTACATTGCTGATTGTGCCTTCCGGGGGCTTTC	
P_AAZ52206	901	ACTGCACATTGGGAACAACAGAGTCAGCTACATTGCTGATTGTGCCTTCCGGGGGCTTTC	
DNA37140	1323	CAGTTTAAAGACTTTGGATCTGAAGAACAATGAAATTTCCTGGACTATTGAAGACATGAA	
P_AAZ52206	961	CAGTTTAAAGACTTTGGATCTGAAGAACAATGAAATTTCCTGGACTATTGAAGACATGAA	
DNA37140	1383	TGGTGCTTTCTCTGGGCTTGACAAACTGAGGCGACTGATACTCCAAGGAAATCGGATCCG	
P_AAZ52206	1021	TGGTGCTTTCTCTGGGCTTGACAAACTGAGGCGACTGATACTCCAAGGAAATCGGATCCG	
DNA37140	1443	TTCTATTACTAAAAAAGCCTTCACTGGTTTGGATGCATTGGAGCATCTAGACCTGAGTGA	
P_AAZ52206	1081	TTCTATTACTAAAAAAGCCTTCACTGGTTTGGATGCATTGGAGCATCTAGACCTGAGTGA	
DNA37140	1503	CAACGCAATCATGTCTTTACAAGGCAATGCATTTTCACAAATGAAGAAACTGCAACAATT	
P_AAZ52206	1141	CAACGCAATCATGTCTTTACAAGGCAATGCATTTTCACAAATGAAGAAACTGCAACAATT	
DNA37140	1563	GCATTTAAATACATCAAGCCTTTTGTGCGATTGCCAGCTAAAATGGCTCCCACAGTGGGT	
P_AAZ52206	1201	GCATTTAAATACATCAAGCCTTTTGTGCGATTGCCAGCTAAAATGGCTCCCACAGTGGGT	
DNA37140	1623	GGCGGAAAACAACTTTCAGAGCTTTGTAAATGCCAGTTGTGCCCATCCTCAGCTGCTAAA	
P_AAZ52206	1261	GGCGGAAAACAACTTTCAGAGCTTTGTAAATGCCAGTTGTGCCCATCCTCAGCTGCTAAA	
DNA37140	1683	AGGAAGAAGCATTTTTGCTGTTAGCCCCAGATGGCTTTGTGTGTG	
P_AAZ52206	1321	AGGAAGAAGCATTTTTGCTGTTAGCCCAGATGGCTTTGTGTGTG	
DNA37140	1743	CCAGATCACGGTTCAGCCAGAAACACAGTCGGCAATAAAAGGTTCCAATTTGAGTTTCAT	
P_AAZ52206	1381	CCAGATCACGGTTCAGCCAGAAACACAGTCGGCAATAAAAGGTTCCAATTTGAGTTTCAT	
DNA37140	1803	CTGCTCAGCTGCCAGCAGCAGTGATTCCCCAATGACTTTTGCTTGGAAAAAAGACAATGA	
P_AAZ52206	1441	CTGCTCAGCTGCCAGCAGCAGTGATTCCCCCAATGACTTTTGCTTGGAAAAAAAGACAATGA	
DNA37140	1863	ACTACTGCATGATGCTGAAATGGAAAATTATGCACACCTCCGGGCCCAAGGTGGCGAGGT	
P_AAZ52206	1501	ACTACTGCATGATGCTGAAATGGAAAATTATGCACACCTCCGGGCCCAAGGTGGCGAGGT	

DNA37140	1923	GATGGAGTATACCACCATCCTTCGGCTGCGCGAGGTGGAATTTGCCAGTGAGGGGAAATA
P_AAZ52206	1561	GATGGAGTATACCACCATCCTTCGGCTGCGCGAGGTGGAATTTGCCAGTGAGGGGAAATA
DNA37140	1983	TCAGTGTGTCATCTCCAATCACTTTGGTTCATCCTACTCTGTCAAAGCCAAGCTTACAGT
P_AAZ52206	1621	TCAGTGTCATCTCCAATCACTTTGGTTCATCCTACTCTGTCAAAGCCAAGCTTACAGT
DNA37140	2043	AAATATGCTTCCCTCATTCACCAAGACCCCCATGGATCTCACCATCCGAGCTGGGGCCAT ***********************************
P_AAZ52206	1681	AAATATGCTTCCCTCATTCACCAAGACCCCCATGGATCTCACCATCCGAGCTGGGGCCAT
DNA37140	2103	GGCACGCTTGGAGTGTGCTGCTGTGGGGCACCCAGCCCCCCAGATAGCCTGGCAGAAGGA
P_AAZ52206	1741	GGCACGCTTGGAGTGTGCTGTGGGGCACCCAGCCCCCCAGATAGCCTGGCAGAAGGA
DNA37140	2163	TGGGGGCACAGACTTCCCAGCTGCACGGGAGAGACGCATGCAT
P_AAZ52206	1801	TGGGGGCACAGACTTCCCAGCTGCACGGGAGAGACGCATGCAT
DNA37140	2223	CGTGTTCTTTATCGTGGATGTGAAGATAGAGGACATTGGGGTATACAGCTGCACAGCTCA
P_AAZ52206	1861	$\tt CGTGTTCTTTATCGTGGATGTGAAGATAGAGGACATTGGGGTATACAGCTGCACAGCTCA$
DNA37140	2283	GAACAGTGCAGGAAGTATTTCAGCAAATGCAACTCTGACTGTCCTAGAAACACCATCATT
P_AAZ52206	1921	GAACAGTGCAGGAAGTATTTCAGCAAATGCAACTCTGACTGTCCTAGAAACACCATCATT
DNA37140	2343	TTTGCGGCCACTGTTGGACCGAACTGTAACCAAGGGAGAAACAGCCGTCCTACAGTGCAT
P_AAZ52206	1981	TTTGCGGCCACTGTTGGACCGAACTGTAACCAAGGGAGAAACAGCCGTCCTACAGTGCAT
DNA37140	2403	TGCTGGAGGAAGCCCTCCCCCTAAACTGAACTGGACCAAAGATGATAGCCCATTGGTGGT
P_AAZ52206	2041	TGCTGGAGGAAGCCCTCCCCCTAAACTGAACTGGACCAAAGATGATAGCCCATTGGTGGT
DNA37140	2463	AACCGAGAGGCACTTTTTTGCAGCAGGCAATCAGCTTCTGATTATTGTGGACTCAGATGT
P_AAZ52206	2101	AACCGAGAGGCACTTTTTTGCAGCAGGCAATCAGCTTCTGATTATTGTGGACTCAGATGT
DNA37140	2523	CAGTGATGCTGGGAAATACACATGTGAGATGTCTAACACCCTTGGCACTGAGAGAGGAAA
P_AAZ52206	2161	CAGTGATGCTGGGAAATACACATGTGAGATGTCTAACACCCTTGGCACTGAGAGAGA
DNA37140	2583	CGTGCGCCTCAGTGTGATCCCCACTCCAACCTGCGACTCCCCTCAGATGACAGCCCCATC
P_AAZ52206	2221	CGTGCGCCTCAGTGTGATCCCCACTCCAACCTGCGACTCCCCTCAGATGACAGCCCCATC
DNA37140	2643	GTTAGACGATGACGGATGGGCCACTGTGGGTGTCGTGATCATAGCCGTGGTTTGCTGTGT **********************
P_AAZ52206	2281	GTTAGACGATGACGGATGGGCCACTGTGGGTGTCGTGATCATAGCCGTGGTTTGCTGTGT
DNA37140	2703	GGTGGGCACGTCACTCGTGTGGGTGGTCATCATATACCACACAAGGCGGAGGAATGAAGA
P_AAZ52206	2341	GGTGGGCACGTCACTCGTGTGGGTGGTCATCATATACCACACAAGGCGGAGGAATGAAGA

DNA37140	2763	TTGCAGCATTACCAACACAGATGAGACCAACTTGCCAGCAGATATTCCTAGTTATTTGTC *********************************
P_AAZ52206	2401	TTGCAGCATTACCAACACAGATGAGACCAACTTGCCAGCAGATATTCCTAGTTATTTGTC
DNA37140	2823	ATCTCAGGGAACGTTAGCTGACAGGCAGGATGGGTACGTGTCTTCAGAAAGTGGAAGCCA *******************************
P_AAZ52206	2461	ATCTCAGGGAACGTTAGCTGACAGGCAGGATGGGTACGTGTCTTCAGAAAGTGGAAGCCA
DNA37140	2883	CCACCAGTTTGTCACATCTTCAGGTGCTGGATTTTTCTTACCACAACATGACAGTAGTGG *******************************
P_AAZ52206	2521	CCACCAGTTTGTCACATCTTCAGGTGCTGGATTTTTCTTACCACAACATGACAGTAGTGG
DNA37140	2943	GACCTGCCATATTGACAATAGCAGTGAAGCTGATGTGGAAGCTGCCACAGATCTGTTCCT ******************************
P_AAZ52206	2581	GACCTGCCATATTGACAATAGCAGTGAAGCTGATGTGGAAGCTGCCACAGATCTGTTCCT
DNA37140		TTGTCCGTTTTTGGGATCCACAGGCCCTATGTATTTGAAGGGAAATGTGTATGGCTCAGA ***********************************
P_AAZ52206		TTGTCCGTTTTTGGGATCCACAGGCCCTATGTATTTGAAGGGAAATGTGTATGGCTCAGA
DNA37140	3063	TCCTTTTGAAACATATCATACAGGTTGCAGTCCTGACCCAAGAACAGTTTTAATGGACCA *********************************
P_AAZ52206	2701	TCCTTTTGAAACATATCATACAGGTTGCAGTCCTGACCCAAGAACAGTTTTAATGGACCA
DNA37140	3123	CTATGAGCCCAGTTACATAAAGAAAAAGGAGTGCTACCCATGTTCTCATCCTTCAGAAGA ********************************
P_AAZ52206	2761	CTATGAGCCCAGTTACATAAAGAAAAAGGAGTGCTACCCATGTTCTCATCCTTCAGAAGA
DNA37140		ATCCTGCGAACGGAGCTTCAGTAATATCGTGGCCTTCACATGTGAGGAAGCTACTTAA *******************************
P_AAZ52206		ATCCTGCGAACGGAGCTTCAGŢAATATATCGTGGCCTTCACATGTGAGGAAGCTACTTAA
DNA37140	3243	CACTAGTTACTCTCACAATGAAGGACCTGGAATGAAAAATCTGTGTCTAAACAAGTCCTC *********************************
P_AAZ52206	2881	CACTAGTTACTCTCACAATGAAGGACCTGGAATGAAAAATCTGTGTCTAAACAAGTCCTC
DNA37140		TTTAGATTTTAGTGCAAATCCAGAGCCAGCGTCGGTTGCCTCGAGTAATTCTTTCATGGG **********************************
P_AAZ52206	2941	TTTAGATTTTAGTGCAAATCCAGAGCCAGCGTCGGTTGCCTCGAGTAATTCTTTCATGGG
DNA37140	3363	TACCTTTGGAAAAGCTCTCAGGAGACCTCACCTAGATGCCTATTCAAGCTTTGGACAGCC *******************************
P_AAZ52206	3001	TACCTTTGGAAAAGCTCTCAGGAGACCTCACCTAGATGCCTATTCAAGCTTTGGACAGCC
DNA37140	3423	ATCAGATTGTCAGCCAAGAGCCTTTTATTTGAAAGCTCATTCTTCCCCAGACTTGGACTC ***********************************
P_AAZ52206	3061	ATCAGATTGTCAGCCAAGAGCCTTTTATTTGAAAGCTCATTCTTCCCCAGACTTGGACTC
DNA37140		TGGGTCAGAGGAAGATGGGAAAGAAAGGACAGATTTTCAGGAAGAAATCACATTTGTAC ************************************
P_AAZ52206		TGGGTCAGAGGAAGAAGAAGAAGGACAGATTTTCAGGAAGAAAATCACATTTGTAC
DNA37140		CTTTAAACAGACTTTAGAAAACTACAGGACTCCAAATTTTCAGTCTTATGACTTGGACAC *********************************
P_AAZ52206		CTTTAAACAGACTTTAGAAAACTACAGGACTCCAAATTTTCAGTCTTATGACTTGGACAC
DNA37140	3603	${\tt ATAGACTGAATGAGACCAAAGGAAAAGCTTAACATACTACCTCAAGTGAACTTTTATTTA$

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         P AAZ52206
         3663 AAAGAGAGAATCTTATGTTTTTTAAATGGAGTTATGAATTTTAAAAGGATAAAAATGC
 DNA37140
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         3301 AAAGAGAGAATCTTATGTTTTTTAAATGGAGTTATGAATTTTAAAAGGATAAAAATGC
P AAZ52206
         3723 TTTATTTATACAGATGAACCAAAATTACAAAAAGTTATGAAAAATTTTTATACTGGGAATG
 DNA37140
             ************
         3361 TTTATTTATACAGATGAACCAAAATTACAAAAAGTTATGAAAATTTTTATACTGGGAATG
P AAZ52206
         3783 ATGCTCATATAAGAATACCTTTTTAAACTATTTTTTAACTTTGTTTTATGCAAAAAAGTA
 DNA37140
             ***********
         3421 ATGCTCATATAAGAATACCTTTTTAAACTATTTTTTAACTTTGTTTTATGCAAAAAAGTA
P AAZ52206
         3843 TCTTACGTAAATTAATGATATAAATCATGATTATTTTATGTATTTTTATAATGCCAGATT
 DNA 37140
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         3481 TCTTACGTAAATTAATGATATAAATCATGATTATTTTATGTATTTTTATAATGCCAGATT
P AAZ52206
         3903 TCTTTTTATGGAAAATGAGTTACTAAAGCATTTTAAATAATACCTGCCTTGTACCATTTT
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 DNA37140
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         3601 TTAAATAGAAGTTACTTCATTATATTTTGCACATTATATTTAATAAAATGTGTCAATTTG
P AAZ52206
        4023 AA
 DNA37140
P AAZ52206
        3661 AA
>10 P AAX52264 Protein PRO335 cDNA clone DNA41388-1234. DNA, PAT 25-JUN-1999
(3662 bp) [1 seg]
Score = 3542 (7022 bits), Expect = 0.0
Identities = 3542/3542 (100%), at 483,121-4024,3662, Strand +/+
          483 GGACTTAAGTCACAACAGATTATCTTTCATCAAGGCAAGTTCCATGAGCCACCTTCAAAG
 DNA37140
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          121 GGACTTAAGTCACAACAGATTATCTTTCATCAAGGCAAGTTCCATGAGCCACCTTCAAAG
P AAX52264
          543 CCTTCGAGAAGTGAAACTGAACAATGAATTGGAGACCATTCCAAATCTGGGACCAGT
 DNA37140
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          181 CCTTCGAGAAGTGAAACTGAACAACAATGAATTGGAGACCATTCCAAATCTGGGACCAGT
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P AAX52264
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 DNA37140
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          361 GCTCCAAACTGCATTTCCAGCCCTACAGCTCAAATATCTGTATCTCAACAGCAACCGAGT
P AAX52264
          783 CACATCAATGGAACCTGGGTATTTTGACAATTTGGCCAACACACTCCTTGTGTTAAAGCT
 DNA37140
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P_AAX52264	421	CACATCAATGGAACCTGGGTATTTTGACAATTTGGCCAACACACTCCTTGTGTTAAAGCT
DNA37140	843	GAACAGGAACCGAATCTCAGCTATCCCACCCAAGATGTTTAAACTGCCCCAACTGCAACA
P_AAX52264	481	
DNA37140	903	TCTCGAATTGAACCGAAACAAGATTAAAAATGTAGATGGACTGACATTCCAAGGCCTTGG
P_AAX52264	541	TCTCGAATTGAACCGAAACAAGATTAAAAATGTAGATGGACTGACATTCCAAGGCCTTGG
DNA37140	963	TGCTCTGAAGTCTCTGAAAATGCAAAGAAATGGAGTAACGAAACTTATGGATGG
P_AAX52264	601	TGCTCTGAAGTCTCTGAAAATGCAAAGAAATGGAGTAACGAAACTTATGGATGG
DNA37140	1023	TTGGGGGCTGAGCAACATGGAAATTTTGCAGCTGGACCATAACAACCTAACAGAGATTAC
P_AAX52264	661	TTGGGGGCTGAGCAACATGGAAATTTTGCAGCTGGACCATAACAACCTAACAGAGATTAC
DNA37140	1083	CAAAGGCTGGCTTTACGGCTTGCTGATGCTGCAGGAACTTCATCTCAGCCAAAATGCCAT
P_AAX52264	721	CAAAGGCTGGCTTTACGGCTTGCTGATGCTGCAGGAACTTCATCTCAGCCAAAATGCCAT
DNA37140	1143	CAACAGGATCAGCCCTGATGCCTGGGAGTTCTGCCAGAAGCTCAGTGAGCTGACCTAAC ********************************
P_AAX52264	781	CAACAGGATCAGCCCTGATGCCTGGGAGTTCTGCCAGAAGCTCAGTGAGCTGACCTAAC
DNA37140	1203	TTTCAATCACTTATCAAGGTTAGATGATTCAAGCTTCCTTGGCCTAAGCTTACTAAATAC ***************************
P_AAX52264	841	TTTCAATCACTTATCAAGGTTAGATGATTCAAGCTTCCTTGGCCTAAGCTTACTAAATAC
DNA37140	1263	ACTGCACATTGGGAACAACAGÅGTCAGCTACATTGCTGATTGTGCCTTCCGGGGGCTTTC ******************
P_AAX52264	901	ACTGCACATTGGGAACAACAGAGTCAGCTACATTGCTGATTGTGCCTTCCGGGGGCTTTC
DNA37140	1323	CAGTTTAAAGACTTTGGATCTGAAGAACAATGAAATTTCCTGGACTATTGAAGACATGAA
P_AAX52264	961	CAGTTTAAAGACTTTGGATCTGAAGAACAATGAAATTTCCTGGACTATTGAAGACATGAA
DNA37140	1383	TGGTGCTTTCTCTGGGCTTGACAAACTGAGGCGACTGATACTCCAAGGAAATCGGATCCG ***********************************
P_AAX52264	1021	TGGTGCTTTCTCTGGGCTTGACAAACTGAGGCGACTGATACTCCAAGGAAATCGGATCCG
DNA37140	1443	TTCTATTACTAAAAAAGCCTTCACTGGTTTGGATGCATTGGAGCATCTAGACCTGAGTGA ********************************
P_AAX52264	1081	TTCTATTACTAAAAAAGCCTTCACTGGTTTGGATGCATTGGAGCATCTAGACCTGAGTGA
DNA37140	1503	CAACGCAATCATGTCTTTACAAGGCAATGCATTTTCACAAATGAAGAAACTGCAACAATT *******************************
P_AAX52264	1141	CAACGCAATCATGTCTTTACAAGGCAATGCATTTTCACAAATGAAGAAACTGCAACAATT
DNA37140	1563	GCATTTAAATACATCAAGCCTTTTGTGCGATTGCCAGCTAAAATGGCTCCCACAGTGGGT ********************************
P_AAX52264	1201	GCATTTAAATACATCAAGCCTTTTGTGCGATTGCCAGCTAAAATGGCTCCCACAGTGGGT
DNA37140	1623	GGCGGAAAACAACTTTCAGAGCTTTGTAAATGCCAGTTGTGCCCATCCTCAGCTGCTAAA **********************************

P_AAX52264	1261	GGCGGAAAACAACTTTCAGAGCTTTGTAAATGCCAGTTGTGCCCATCCTCAGCTGCTAAA
DNA37140	1683	AGGAAGAAGCATTTTTGCTGTTAGCCCAGATGGCTTTGTGTGTG
P_AAX52264	1321	AGGAAGAAGCATTTTTGCTGTTAGCCCAGATGGCTTTGTGTGTG
DNA37140	1743	CCAGATCACGGTTCAGCCAGAAACACAGTCGGCAATAAAAGGTTCCAATTTGAGTTTCAT
P_AAX52264	1381	${\tt CCAGATCACGGTTCAGCCAGAAACACAGTCGGCAATAAAAGGTTCCAATTTGAGTTTCAT}$
DNA37140	1803	CTGCTCAGCTGCCAGCAGCAGTGATTCCCCCAATGACTTTTGCTTGGAAAAAAGACAATGA **********************************
P_AAX52264	1441	CTGCTCAGCTGCCAGCAGCAGTGATTCCCCCAATGACTTTTGCTTGGAAAAAAGACAATGA
DNA37140	1863	ACTACTGCATGATGCTGAAATGGAAAATTATGCACACCTCCGGGCCCAAGGTGGCGAGGT
P_AAX52264	1501	ACTACTGCATGATGCTGAAATGGAAAATTATGCACACCTCCGGGCCCAAGGTGGCGAGGT
DNA37140	1923	GATGGAGTATACCACCATCCTTCGGCTGCGCGAGGTGGAATTTGCCAGTGAGGGGAAATA *************************
P_AAX52264	1561	GATGGAGTATACCACCATCCTTCGGCTGCGCGAGGTGGAATTTGCCAGTGAGGGGAAATA
DNA37140	1983	TCAGTGTGTCATCTCCAATCACTTTGGTTCATCCTACTCTGTCAAAGCCAAGCTTACAGT ************************************
P_AAX52264	1621	TCAGTGTGTCATCTCCAATCACTTTGGTTCATCCTACTCTGTCAAAGCCAAGCTTACAGT
DNA37140	2043	AAATATGCTTCCCTCATTCACCAAGACCCCCATGGATCTCACCATCCGAGCTGGGGCCAT ***********************************
P_AAX52264	1681	AAATATGCTTCCCTCATTCACCAAGACCCCCATGGATCTCACCATCCGAGCTGGGGCCAT
DNA37140	2103	GGCACGCTTGGAGTGTGCTGCTGTGGGGCACCCAGCCCCCCAGATAGCCTGGCAGAAGGA *****************************
P_AAX52264	1741	GGCACGCTTGGAGTGTGCTGTGGGGCACCCAGCCCCCAGATAGCCTGGCAGAAGGA
DNA37140	2163	TGGGGGCACAGACTTCCCAGCTGCACGGGAGAGACGCATGCAT
P_AAX52264	1801	TGGGGGCACAGACTTCCCAGCTGCACGGGAGAGACGCATGCAT
DNA37140	2223	CGTGTTCTTTATCGTGGATGTGAAGATAGAGGACATTGGGGTATACAGCTGCACAGCTCA ***********************************
P_AAX52264	1861	CGTGTTCTTTATCGTGGATGTGAAGATAGAGGACATTGGGGTATACAGCTGCACAGCTCA
DNA37140	2283	GAACAGTGCAGGAAGTATTTCAGCAAATGCAACTCTGACTGTCCTAGAAACACCATCATT **************************
P_AAX52264	1921	GAACAGTGCAGGAAGTATTTCAGCAAATGCAACTCTGACTGTCCTAGAAACACCATCATT
DNA37140	2343	TTTGCGGCCACTGTTGGACCGAACTGTAACCAAGGGAGAAACAGCCGTCCTACAGTGCAT ************************************
P_AAX52264	1981	TTTGCGGCCACTGTTGGACCGAACTGTAACCAAGGGAGAAACAGCCGTCCTACAGTGCAT
DNA37140	2403	TGCTGGAGGAAGCCCTCCCCCTAAACTGAACTGGACCAAAGATGATAGCCCATTGGTGGT
P_AAX52264	2041	TGCTGGAGGAAGCCCTCCCCCTAAACTGAACTGGACCAAAGATGATAGCCCATTGGTGGT
DNA37140	2463	AACCGAGAGGCACTTTTTTGCAGCAGGCAATCAGCTTCTGATTATTGTGGACTCAGATGT *********************************
P_AAX52264	2101	AACCGAGAGGCACTTTTTTGCAGCAGGCAATCAGCTTCTGATTATTGTGGACTCAGATGT

DNA37140 2523 CAGTGATGCTGGGAAATACACATGTGAGATGTCTAACACCCTTGGCACTGAGAGGGAAA  P_AAX52264 2161 CAGTGATGCTGGGAAATACACACTGTAACACCCTTGGCACTGAGAGGGAAA  DNA37140 2593 CGTGCGCCTCAGTGTGATCCCCACTCCAACCTGCGACTCCCCCTCAGATGACAGCCCATC  P_AAX52264 2221 CGTGCGCCTCAGTGTGATCCCCACTCCAACCTGCGACTCCCCCTCAGATGACAGCCCCATC  DNA37140 2643 CTTAGACGATGACGCATGGGCCACTGTGGGTGTCGTGATCATACCCGTTGGTTTGCTGTGT  P_AAX52264 2281 GTTAGACGATGACCGATGGGCCACTGTGGGTGTCGTGATCATAGCCGTGGTTTGCTGTGT  DNA37140 2703 GGTGGCACCTCACTCGTGGGGTGTCATCATATACCACACAAGGCCGGAGGAATGAAGA  DNA37140 2763 TTGCAGCACTCACTCGTGTGGGTGTCATCATATACCACACAAGGCCGGAGGAATGAAGA  DNA37140 2763 TTGCAGCATTACCAACACAGAATGAGCCAACTTGCCAGCAGAGATCTCTTATTTTCTC  P_AAX52264 2401 TTGCAGCATTACCAACACAGATGAGCCAGATTTCCTAGATATTCCTAGTTATTTTCTC  DNA37140 2823 ATCTCAGGGAACCTTACCTGACAGGCCGGAGGTGGTTCTTCAGAAAGTGGAAGCCA  DNA37140 2883 ATCTCAGGGAACCTTACCTGACAGGCAGGATGGGTACGTGTTTCAGAAAGTGGAAGCCA  DNA37140 2883 CCACCAGTTTGCCACACTCTGAGGCAGGATGGGTACGTGTCTTCAGAAAGTGGAAGCCA  DNA37140 2983 GACCTCCCATATTGCAACTCTCACGGCAGGATGTTTTCTTACCACACACA		
P_AAX52264 2161 CAGTGATGCTGGAAATACACATGTGAGATGCTCAGCACTGAGAGAGA		
P_AAX52264         2161         CAGTGATGCTGGGAAATACACATGTGGGATGTCTAACACCCTTGCACTGGAGAGGAGACACTGAAATACACATGTGAGATGTCCCACTCCAACCTGCGACTCCACCTGGATGACAGCCCCATCCAACCTGCAACCTGCACTGAGATGACAGCCCCATCCAACCTGCAACCTGCGACTCCCCACTCCAACCTGCACTCCACACCTGCACTCCACATGACAGACCCCATCCACACTGCACTCCACACCTCCACACCTGCGACTCCCCCACTCCACACCTGCGACTCCCCCACTCCACACCTGCGACTCCCCCACTCGATGACACACCCCCATCCACACCTGCGACTCCCCCCACTCGTGTGTCTCTGTGTTCTCTGTTTCTCTGTTTCTCACACACCACCACCACCTGGATGACCACCACCACCACCGCGAGACACCACCTCGTGTGGGTGCTCACTCA		A37140
P_AAX52264 2221 CGTGCCCTCAGTGTGTGTCCCCACTCCAACCTGCGACTCCCCTCAGATGACAGCCCCATC DNA37140 2643 GTTAGACGATGACGGTGGCCACTGTGGGTGTCGTGATCATAGCCGTGGTTTGCTGTGT  P_AAX52264 2281 GTTAGACGATGACCGATGGGCCACTGTGGGTGTCGTGATCATAGCCGTGGTTTGCTGTGT DNA37140 2703 GGTGGGCACGTCACTCGTGTGGGTGGTCATCATAACCACACAAGGGGGGAGATGAAGA  P_AAX52264 2341 GGTGGGCACGTCACTCGTGTGGGTGGTCATCATATACCACACAAAGGCGGAGGATGAAGA  DNA37140 2763 TTGCAGCATTACCAACACAGATGAGCCACACTTGCCGAGCAGATATTCCTAGTTATTTGTC  DNA37140 2823 ATCTCAGGGAACGTTAGCAACACAGATGAGCCAACTTGCCAGCAGATATTCCTAGTTATTTGTC  DNA37140 2823 ATCTCAGGGAACGTTAGCTGACAGGCAGGATGGGTACGTGTCTTCAGAAAATGGGAAGCCA  DNA37140 2883 CCACCAGTTTGTCACATCTTCAGGTAGGTGGTTACTTACCAACACACAGATGGGAAGCCA  DNA37140 2883 CCACCAGTTTGTCACATCTTCAGGTGCTGGATTTTCTTACCACAACATGACAGTAGTGG  DNA37140 2943 GACCTGCCATATTGACAATAGCAGTGAGCTGGATTTTCTTACCACAACATGACAGTAGTGG  DNA37140 2943 GACCTGCCATATTGACAATAGCAGTGAAGCTGGAAGCTGCCACAGATCTGTTCCT  P_AAX52264 251 GACCTCCCATATTGACAATAGCAGTGAAGCTGGATGTGGAAGCTGCCACAGATCTGTTCCT  DNA37140 3003 TTGTCCGTTTTTGGGATCCACAGGCCCTATGTATTTGAAGGGAAACTGTATGGTCCAGA  P_AAX52264 2641 TTGTCCGTTTTTGGGATCCACAGGCCCTATGTATTTGAAGGGAAACTGTATGGCTCAGA  DNA37140 3003 TCCTTTTGAAACATATCATACAGGTTGCAGTCCTGACCCAAGAACAGTTTTAATGGACCA  DNA37140 3123 CTATGAGCCCAGTTACTAACAGGTTGCAGTCCTGACCCAAGAACAGTTTTAATGGACCA  DNA37140 3123 CTATGAGCCCAGTTACATAAAGAAAAAGGAGTGCTCCACCAGGAACAGTTTTAATGGACCA  DNA37140 3183 ATCCTGCGAACGGAGCTTCAGTAATATATCAGGCCCTATGTACCCATGTTCCTCACATGTCCTTCAGAAGAA  DNA37140 3183 ATCCTGCGAACGGAGCTTCAGTAATATATCTGGCCCTACCATGTTCCTCACATGTAGAGAAAAGAATATCATACAGGTCGAACAGATGTTCACACATGTACAGAGACAGTTTTAAAAAAAA		X52264
P_AAX52264 2221 CGTGCGCTCAGTGTGATCCCCACTCCAACCTGCGACTCCCCTCAGATGACAGCCCCATCTCAGATGACAGCCCCATCTCAGATGACAGCCCCATCTTCAGATGACAGCCCCATCTTCAGATGACAGATGACGATGACGATGACGATGACGATGACGATGACGATGACGATGACGATGACGATGACGATGACGATGACGATGACGATGACGATGACCACAAAGCCGGAGAATGAAGA  P_AAX52264 2341 GGTGGGCACGTCACTCGTGTGGGTGGTCATCATATACCACACAAGGCCGAGGAATGAAGA  DNA37140 2763 TTGCAGCATTACCAACACAGAGAGCAGACTTAGCCACAAAGCCGAGAGAATTATCTC  P_AAX52264 2401 TTGCAGCATTACCAACACAGAGAGCAGACTTGCCAGCAGAGATATTCCTAGTTATTTGTC  DNA37140 2823 ATCTCAGGGAACGTTAGCTGACAGGCAGGATGGGTACGTGTCTTCAGAAAGTGGAAGCCA  DNA37140 2883 CCACCAGTTTGTCACATCTTCAGGAGGCAGGATGGTTCCTTCAGAAAGTGGAAGCCA  DNA37140 2883 CCACCAGTTTGTCACATCTTCAGGTGCTGGATTTTTCTTACCACACACA		A37140
P_AAX52264 2281 GTTAGACGATGACGGATGGGCCACTGTGGGTTCGTGATCATAGCCGTGGTTTGCTGTGT DNA37140 2703 GGTGGGCACGTCACTCGTGTGGTTGGTTGATCATATACCACACAAGGCGAGGAATGAAGA P_AAX52264 2341 GGTGGGCACGTCACTCGTGTGGGTGGTCATCATATACCACACAAGGCGAGGAATGAAGA DNA37140 2763 TTGCAGCATTACCAACACAGATGAGACCAACTTGCCAGCAGATATTCCTAGTTATTTGTC DNA37140 2823 ATCTCAGGGAACGTTAGCTGACAGGCAGGATGGTACGTTCTTCAGAAAGTGGAAGCCA DNA37140 2883 ACCTCAGGGAACGTTAGCTGACAGGCAGGATGGTACGTTCTTCAGAAAGTGGAAGCCA DNA37140 2883 CCACCAGTTTGTCACATCTTCAGGATTGTTTTTTTTCTCAAACACACAAGATGAGACCAACATGGCAGGATGGTACGTGTCTTCAGAAAGTGGAAGCCA DNA37140 2883 CCACCAGTTTGTCACATCTTCAGGTGCTGGATTTTTCTTACCACAACATGACAGTAGTGG DNA37140 2943 GACCTGCCATATTGACAATAGCAGTGAGAGCTGATTTTCTTACCACAACATGACAGTAGTGG DNA37140 3003 TTGTCCGTTTTTGGGATCCACAGGCCGTGATGTTGGAAGCTGCACAGATCTGTTCCT DNA37140 3003 TTGTCCGTTTTTGGGATCCACAGGCCCTATGTATTTGAAGGGAAACTGTTTAGTGTCCAGA DNA37140 3003 TCCTTTTGGAACATACAATACAGGTGCACCAAGAACAGTTTTAATGGACCA DNA37140 3063 TCCTTTTGGAACATATCATACAGGTGCAGTCCTGACCCAAGAACAGTTTTAATGGACCA DNA37140 3123 CTATGAGCCCAGTTACATAAAGAAAAAGGAGTGCTACCCAAGAACAGTTTTAATGGACCA DNA37140 3123 CTATGAGCCCAGTTACATAAAGAAAAAGGAGTGCTACCCAAGAACAGTTTTAATGGACCA DNA37140 3123 CTATGAGCCCAGTTACATAAAGAAAAAAGGAGTGCTACCCATGTTCCATCCTTCAGAAGA DNA37140 3123 CTATGAGCCCAGTTACATAAAGAAAAAAGGAGTGCTACCCATGTTCCATCCTTCAGAAGA DNA37140 3123 CTATGAGCCCAGTTACATAAAGAAAAAAGGAGTGCTACCCATGTTCCATCCTTCAGAAGA DNA37140 3123 CTATGAGCCCAGTTACATAAAGAAAAAAGGAGTGCTACCCATGTTCCATCCTTCAGAAGA DNA37140 3123 CTATGAGCCCAGTTACATAAAGAAAAAAGGAGTGCTACCCATGTTCACATGTGAGGAAGCTACTTAA DNA37140 3123 CACTAGTTACTCCACAATGAAGAACAGTCTGAGAACAAACA		X52264
DNA37140 2703 GGTGGGCACGTCACTCGTGTGGGTGGTCATCATATACCACACAAGGCGGAGGAATGAAGA P_AAX52264 2341 GGTGGGCACGTCACTCGTGTGGGTGGTCATCATATACCACACAAGGCGGAGGAATGAAGA DNA37140 2763 TTGCAGCATTACCAACACAGATGAGCCAACATGCCAGCAGATATTCCTAGTTATTTGTC P_AAX52264 2401 TTGCAGCATTACCAACACAGATGAGACCAACTTGCCAGCAGATATTCCTAGTTATTTGTC DNA37140 2823 ATCTCAGGGAACGTTAGCTGACAGGCAGGATGGGTACGTGTCTTCAGAAAGTGGAAGCCA DNA37140 2883 CCACCAGTTTGTCACATCTTCAGGTGGCTGGCATGTTCTTCAGAAAGTGGAAGCCA DNA37140 2883 CCACCAGTTTGTCACATCTTCAGGTGCTGGATTTTTCTTACCACACACA		A37140
P_AAX52264 2341 GGTGGGCACGTCACTCGTGTGGGTGGTCATCATATACCACACAGAGGCGGAGGAATGAAGA  DNA37140 2763 TTGCAGCATTACCAACACAGAGAGAGACCAACTTGCCAGCAGAGATATTCCTAGTTATTTGTC  P_AAX52264 2401 TTGCAGCATTACCAACACAGAGAGAGACCAACTTGCCAGCAGATATTCCTAGTTATTTTGTC  DNA37140 2823 ATCTCAGGGAACGTTAGCTGACAGGCAGGATGGGTACGTGTCTTCAGAAAGTGGAAGCCA  P_AAX52264 2461 ATCTCAGGGAACGTTAGCTGACAGGCAGGATGGGTACGTGTCTTCAGAAAGTGGAAGCCA  DNA37140 2883 CCACCAGTTTGTCACATCTTCAGGTGCTGGATTTTTCTTACCACAACATGACAGTAGTGG  P_AAX52264 2521 CCACCAGTTTGTCACATCTTCAGGTGCTGGATTTTTCTTACCACAACATGACAGTAGTGG  DNA37140 2943 GACCTGCCATATTGACAATAGCAGTGAGGCTGATGTGGAAGCTGCCACAGATCTGTTCCT  DNA37140 3003 TTGTCCGTTTTTGGGATCCACAGGCCGATGTGTGGAAGCTGCCACAGATCTGTTCCT  DNA37140 3063 TCCTTTTGGAACATACCAAGGCCCTATGTATTTGAAGGGAAATGTTATTGGCTCAGA  DNA37140 3063 TCCTTTTGAAACATATCATACAGGTTGCAGTCCTGACCCAAGAACAGTTTTAATGGACCA  DNA37140 3123 CTATGAGCCCAGTTACATAAAGAAAAAGGAGTGCTACCCAAGAACAGTTTTAATGGACCA  DNA37140 3123 CTATGAGCCCAGTTACATAAAGAAAAAGGAGTGCTACCCATGTTCTCATCCTTCAGAAGA  DNA37140 3183 ATCCTGCGAACGGAGCTTCAGTAATATATCGTGCCCCAAGAACAGTTTTAATGGACCA  DNA37140 3183 ATCCTGCGAACGGAGCTTCAGTAATATATCGTGGCCTTCACATGTTGCTCTCAGAAGA  DNA37140 3183 ATCCTGCGAACGGAGCTTCAGTAATATATTCGTGGCCTTCACATGTGAGGAAGCTACTTAA  DNA37140 3183 CACTAGTTACTCACAATGAAGAAAAAAGGAGTTCTCACATGTGAGGAAGCTACTTAA  DNA37140 3183 CACTAGTTACTTCACAATGAAGAAAAAAGGAGTTCTCACATGTGAGGAAGCTACTTAA  DNA37140 3183 CACTAGTTACTTCACAATGAAGAACAGGCCCTTCACATGTGAGGAAGCTACTTAA  DNA37140 3183 CACTAGTTACTCACAATGAAGAACAGGCCCTGAACAAAAATCTGTGTCTCAACAAGCCCACTATTAACAAGACCCCACGAACAGGACCTACCCAAGAACAGTTCTAAACAAGTCCTC  P_AAX52264 2881 CACTAGTTACTCCACAATGAAGGACCTGGAATGAAAAAATCTGTGTCTAAACAAGTCCTC  P_AAX52264 2881 CACTAGTTACTCTCACAATGAAGGACCTGGAATGAAAAAATCTGTGTCTAAACAAGTCCTC  P_AAX52264 2881 CACTAGTTACTCCACAATGAAGGACCTGGAATGAAAAAATCTGTGTCTAAACAAGTCCTC  DNA37140 330 TTTAGATTTTAGTGCAAATCCAGGGCCGGGTCGGTTGCCTCGAGTAATTCTTTAAACAAGTCCTC  DNA37140 330 TTTAGATTTTAGTGCAAATCCAGGGCCAGGGTCGGTTGCCTCGAGTAATTCTTTCATGGCCTTTAAACAAGTCCTC	_AAX52264 2281 GTTAGACGATGACGGATGGGCCACTGTGGGTGTCGTGATCATAGCCGTGGTTTGCTGTGT	X52264
DNA37140 2763 TTGCAGCATTACCAACACAGATGAGACCAACTTGCCAGCAGATATTCCTAGTTATTTGTC  P_AAX52264 2401 TTGCAGCATTACCAACACAGATGAGACCAACTTGCCAGCAGATATTCCTAGTTATTTGTC  DNA37140 2823 ATCTCAGGGAACGTTAGCTGACAGGCAGGATGGTACGTGTCTTCAGAAAGTGGAAGCCA  P_AAX52264 2461 ATCTCAGGGAACGTTAGCTGACAGGCAGGATGGTACGTGTCTTCAGAAAGTGGAAGCCA  DNA37140 2883 CCACCAGTTTGTCACATCTTCAGGTGCTGGATTTTTCTTACCACAACATGACAGTAGTGG  DNA37140 2943 GACCTGCCATATTGACAATAGCAGTGGAGTTTTTCTTACCACAACATGACAGTAGTGG  DNA37140 2943 GACCTGCCATATTGACAATAGCAGTGAGGTGGTGGAAGCTGCACAGATCTGTTCCT  P_AAX52264 2581 GACCTGCCATATTGACAATAGCAGTGAAGCTGATGTGGAAGCTGCCACAGATCTGTTCCT  DNA37140 3003 TTGTCCGTTTTTGGGATCCACAGGCCCTATGTATTTGAAGGGAAATGTGTATGGCTCAGA  DNA37140 3063 TCCTTTTGGAACCATACCAAGGCCCTATGTATTTGAAGGGAAATGTGTATGGCTCAGA  DNA37140 3063 TCCTTTTGAAACATATCATACAGGTTGCAGTCCTGACCCAAGAACAGTTTTAATGGACCA  DNA37140 3123 CTATGAGCCCAGTTACATAAAGAAAAAGGAGTGCTACCCAAGAACAGTTTTAATGGACCA  DNA37140 3123 CTATGAGCCCAGTTACATAAAGAAAAAAGGAGTGCTACCCATGTTCTCATCCTTCAGAAGA  DNA37140 3183 ATCCTGCGAACGGAGCTTCAGTAATATATCGTGGCCTTCACATGTGAGGAAGCTACTTAA  DNA37140 3183 ATCCTGCGAACGGAGCTTCAGTAATATATCGTGGCCTTCACATGTGAGGAAACAGTTTTAATGAACAAGAC  DNA37140 3183 ATCCTGCGAACGGAGCTTCAGTAATATATATCGTGGCCTTCACATGTGAGGAAACCTACTTAA  DNA37140 3183 CACTAGTTACTCACAATGAAGGACCTGGAATGAAAAATCTGTGTCAGAGAAACACTTCTAA  DNA37140 3183 CACTAGTTACTCACAATGAAGGACCTGGAATGAAAAATCTGTGTCAGAGAAACCTACTTAA  DNA37140 3183 CACTAGTTACTCTCACAATGAAGGACCTGGAATGAAAAATCTGTGTCTAAACAAGCTCCTC  P_AAX52264 2881 CACTAGTTACTCTCACAATGAAGGACCTGGAATGAAAAATCTGTGTCTTAAACAAGTCCTC  P_AAX52264 2881 CACTAGTTACTCTCACAATGAAGGACCTGGAATGAAAAAATCTGTGTCTTAAACAAGTCCTC  DNA37140 3303 TTTAGATTTTAGTGCAAATCCAGGGCCTGGATTGCCTCGAGTAATTCTTTCATCGTGCCAGGAGCCTCTCACATGTTACTCTTCAACAAGTCCTC  DNA37140 3303 TTTAGATTTTAGTGCAAATCCAGGGCCTGGGTTGCCTCGAGTAATTCTTTCATCGTTCTAACCAAGTCCTC  DNA37140 3303 TTTAGATTTTAGTGCAAATCCAGGGCCTGGGTTGCCTCGAGTAATTCTTTCATCGTTCAACAAGTCCTC  DNA37140 3303 TTTAGATTTTAGTGCAAATCCAGAGCCAGGGCTCGGTTGCCTCGAGTAATTCTTTCATCGTGCCTAACCAAGTCCTC		A37140
P_AAX52264 2401 TTGCAGCATTACCAACAGATGAGACCAACTTGCCAGCAGATATTCCTAGTTATTTGTC  DNA37140 2823 ATCTCAGGGAACGTTAGCTGACAGGCAGGATGGGTACGTGTCTTCAGAAAGTGGAAGCCA  P_AAX52264 2461 ATCTCAGGGAACGTTAGCTGACAGGCAGGATGGGTACGTGTCTTCAGAAAGTGGAAGCCA  DNA37140 2883 CCACCAGTTTGTCACATCTTCAGGTGCTGGATTTTTCTTACCACAACATGACAGTAGTGG  P_AAX52264 2521 CCACCAGTTTGTCACATCTTCAGGTGCTGGATTTTTCTTACCACAACAATGACAGTAGTGG  DNA37140 2943 GACCTGCCATATTGACAATAGCAGTGAAGCTGGAAGCTGGCACAGATCTGTTCCT  DNA37140 3003 TTGTCCGTTTTTGGGATCCACAGGCCCTATGTATTTGAAAGGAAATGTGTATGGCTCAGA  DNA37140 3003 TCCTTTTGAAACATATCATACAGGCCCTATGTATTTGAAGGGAAATGTGTATGGCTCAGA  DNA37140 3103 TCCTTTTGAAACATATCATACAGGTTGCAGCCCAAGAACAGTTTTAATGGACCA  DNA37140 3123 CTATGAGCCCAGTTACATAAAGAAAAAGGAGTGCTACCCATGTTCTCATCCTTCAGAAGA  DNA37140 3123 CTATGAGCCCAGTTACATAAAGAAAAAAGGAGTGCTACCCATGTTCTCATCCTTCAGAAGA  DNA37140 3183 ATCCTGCGAACGGAGCTTCAGTAATATATCTGTGGCCTCACATGTAGGAGAACAGTTTTAATGAACA  DNA37140 3183 ATCCTGCGAACGGAGCTTCAGTAATATATCTGTGGCCTTCACATGTAGGAGAACAGTTTTAATGAACATATAA  P_AAX52264 2821 ATCCTGCGAACGGAGCTTCAGTAATATATCTGTGGCCTTCACATGTAGGAAGACTACTTAA  DNA37140 3183 CACTAGTTACTCTCACAATGAAGGACCTGGAATGAAAAATCTGTGTCTAAACAAGTCCTC  P_AAX52264 2881 CACTAGTTACTCTCACAATGAAGGACCTGGAATGAAAAAATCTGTGTCTAAACAAGTCCTC  P_AAX52264 2881 CACTAGTTACTCTCACAATGAAGGACCTGGAATGAAAAAATCTGTGTCTAAACAAGTCCTC  DNA37140 3303 TTTAGATTTTAGTGCAAATCCAGAGCCAGCGTCGGATGGAATAAAATCTGTGTCTAAACAAGTCCTC  DNA37140 3303 TTTAGATTTTAGTGCAAATCCAGAGCCAGCGTCGGTTGCCTCGAGTAAATTCTTTCATGGGCCTAAAACAAGTCCTC  DNA37140 3303 TTTAGATTTTAGTGCAAATCCAGAGCCAGCGTCGGTTGCCTCGAGTAAATTCTTTCATGGGCCTAAAAAAATCTTTTCATCGGGCCTTCAAAAAAAA		X52264
DNA37140 2823 ATCTCAGGGAACGTTAGCTGACAGGCAGGATGGTACGTGTCTTCAGAAAGTGGAAGCCA P_AAX52264 2461 ATCTCAGGGAACGTTAGCTGACAGGCAGGATGGGTACGTGTCTTCAGAAAGTGGAAGCCA DNA37140 2883 CCACCAGTTTGTCACATCTTCAGGTGCTGGATTTTTCTTACCACAACATGACAGTAGTGG P_AAX52264 2521 CCACCAGTTTGTCACATCTTCAGGTGCTGGATTTTTCTTACCACAACATGACAGTAGTGG DNA37140 2943 GACCTGCCATATTGACAATAGCAGTGAGGCTGATGTGGAAGCTGCCACAGATCTGTTCCT DNA37140 3003 TTGTCCGTTTTTGGGATCCACAGGCCCTATGTATTTGAAGGGAAACTGGTATGGCTCAGA DNA37140 3003 TTGTCCGTTTTTGGGATCCACAGGCCCTATGTATTTGAAGGGAAATGTGTATGGCTCAGA DNA37140 3063 TCCTTTTGAAACATATCATACAGGTTGCAGACCCAAGAACAGTTTTAATGGACCA DNA37140 3103 TCCTTTTGAAACATATCATACAGGTTGCAGTCCTGACCCAAGAACAGTTTTAATGGACCA DNA37140 3123 CTATGAGCCCAGTTACATAAAGAAAAAAGGAGTGCTACCCATGTTCTCATCCTTCAGAAGA DNA37140 3123 CTATGAGCCCAGTTACATAAAGAAAAAAGGAGTGCTACCCATGTTCTCATCCTTCAGAAGA DNA37140 3183 ATCCTGCGAACGGAGCTTCAGTAATATATCGTGGCCTTCACATGTGAGGAAGCTACTTAA P_AAX52264 2821 ATCCTGCGAACGGAGCTTCAGTAATATATCGTGGCCTTCACATGTGAGGAAGCTACTTAA DNA37140 3243 CACTAGTTACTCTCACAATGAAGGACCTGGAATGAAAAATCTGTGTCTAAACAAGTCCTC P_AAX52264 2881 CACTAGTTACTCTCACAATGAAGGACCTGGAATGAAAAATCTGTGTCTAAACAAGTCCTC DNA37140 3243 CACTAGTTACTCTCACAATGAAGGACCTGGAATGAAAAATCTGTGTCTAAACAAGTCCTC P_AAX52264 2881 CACTAGTTACTCTCACAATGAAGGACCTGGAATGAAAAATCTGTGTCTAAACAAGTCCTC DNA37140 3303 TTTAGATTTTAGTGCAAATCCAGAGGCCAGCGTCGGTTGCCTCGAGTAATTCTTTCATGGG		A37140
P_AAX52264 2461 ATCTCAGGGAACGTTAGCTGACAGGCAGGATCGTTCTCAGAAAAGTGGAAGCCA  DNA37140 2883 CCACCAGTTTGTCACATCTTCAGGTGCTGGATTTTTCTTACCACAACATGACAGTAGTGG  P_AAX52264 2521 CCACCAGTTTGTCACATCTTCAGGTGCTGGATTTTTCTTACCACAACATGACAGTAGTGG  DNA37140 2943 GACCTGCCATATTGACAATAGCAGTGAAGCTGATGTGGAAGCTGCCACAGATCTGTTCCT  DNA37140 3003 TTGTCCGTTTTTGGGATCCACAGGCCCTATGTATTTGAAGGGAAACTGTGTATCTTCAGAAGA  DNA37140 3063 TCCTTTTGGGATCCACAGGCCCTATGTATTTGAAGGGAAATGTGTATGGCCCACAGATCTGTCCAGAACA  DNA37140 3063 TCCTTTTGAAACATATCATACAGGTTGCAGTCCTGACCCAAGAACAGTTTTAATGGACCA  DNA37140 3123 CTATGAGCCCAGTTACATAAAGAAAAAGGAGTGCTACCCAAGAACAGTTTTAATGGACCA  DNA37140 3123 CTATGAGCCCAGTTACATAAAGAAAAAAGGAGTGCTACCCATGTTCTCATCCTTCAGAAGA  DNA37140 3183 ATCCTGCGAACGGAGCTTCAGTAATATATCCTGGCCTTCACATGTGAGGAAGCTACTTAA  DNA37140 3183 ATCCTGCGAACGGAGCTTCAGTAATATATCTGTGGCCTTCACATGTGAGGAAGCTACTTAA  DNA37140 3243 CACTAGTTACTCTCACAATGAAGGACCTGGAATGAAAAATCTGTGTCTAAACAAGTCCTC  P_AAX52264 2821 ATCCTGCGAACGGAGCTTCAGTAATATATCTGTGGCCTTCACATGTGAGGAAGCTACTTAA  DNA37140 3243 CACTAGTTACTCTCACAATGAAGGACCTGGAATGAAAAATCTGTGTCTAAACAAGTCCTC  P_AAX52264 2881 CACTAGTTACTCTCACAATGAAGGACCTGGAATGAAAAATCTGTGTCTAAACAAGTCCTC  DNA37140 3303 TTTAGATTTTAGTGCAAATCCAGAGCCAGGGTCGGTTGCCTCGAGTAATTTTCATTGGGC  DNA37140 3303 TTTAGATTTTAGTGCAAATCCAGAGCCAGGGTCGGTTGCCTCGAGTAATTTTTTTT	_AAX52264 2401 TTGCAGCATTACCAACACAGATGAGACCAACTTGCCAGCAGATATTCCTAGTTATTTGTC	X52264
DNA37140 2883 CCACCAGTTTGTCACATCTTCAGGTGCTGGATTTTTCTTACCACAACATGACAGTAGTGG P_AAX52264 2521 CCACCAGTTTGTCACATCTTCAGGTGCTGGATTTTTCTTACCACAACATGACAGTAGTGG DNA37140 2943 GACCTGCCATATTGACAATAGCAGTGAAGCTGATGTGGAAGCTGCCACAGATCTGTTCCT P_AAX52264 2581 GACCTGCCATATTGACAATAGCAGTGAAGCTGATGTGGAAGCTGCCACAGATCTGTTCCT DNA37140 3003 TTGTCCGTTTTTGGGATCCACAGGCCCTATGTATTTGAAGGGAAATGTGTATGGCTCAGA DNA37140 3063 TCCTTTTGAAACATATCATACAGGCCCTATGTATTTGAAGGGAAATGTGTATGGCTCAGA DNA37140 3123 CTATTGAAACATATCATACAGGTTGCAGTCCTGACCCAAGAACAGTTTTAATGGACCA DNA37140 3123 CTATGAGCCCAGTTACATAAAGAAAAAGGAGTGCTACCCATGTTCTCATCCTTCAGAAGA DNA37140 3123 CTATGAGCCCAGTTACATAAAGAAAAAAGGAGTGCTACCCATGTTCTCATCCTTCAGAAGA DNA37140 3183 ATCCTGCGAACGGAGCTTCAGTAATATATCGTGGCCTTCACATGTGAGGAAGCTACTTAA DNA37140 3243 CACTAGTTACATCACAATGAAGGACCTGGACTCACATGTGAGGAAGCTACTTAA DNA37140 3243 CACTAGTTACTCTCACAATGAAGGACCTGGAATGAAAAAATCTGTGTCTAAACAAGTCCTC P_AAX52264 2881 CACTAGTTACTCTCACAATGAAGGACCTGGAATGAAAAAATCTGTGTCTAAACAAGTCCTC DNA37140 3243 CACTAGTTACTCTCACAATGAAGGACCTGGAATGAAAAAATCTGTGTCTTAAACAAAGTCCTC DNA37140 3243 CACTAGTTACTCTCACAATGAAGGACCTGGAATGAAAAAATCTGTGTCTTAAACAAAGTCCTC DNA37140 3303 TTTAGATTTTAGTGCCAAATCCAGAGCCAGCGTCGGTTGCCTCGAGTAAATTCTTTCATGGG		A37140
P_AAX52264 2521 CCACCAGTTTGTCACATCTTCAGGTGCTGGATTTTCTTACCACAACATGACAGTAGTGG  DNA37140 2943 GACCTGCCATATTGACAATAGCAGTGAAGCTGGAAGCTGGCCACAGATCTGTTCCT  P_AAX52264 2581 GACCTGCCATATTGACAATAGCAGTGAAGCTGGTAGTGTGGAAGCTGCCACAGATCTGTTCCT  DNA37140 3003 TTGTCCGTTTTTGGGATCCACAGGCCCTATGTATTTGAAGGGAAATGTGTATGGCTCAGA  DNA37140 3063 TCCTTTTGAAACATATCATACAGGTTGCAGTCCTGACCCAAGAACAGTTTTAATGGACCA  DNA37140 3123 CTATGAGCCCAGTTACATACAGGTTGCAGTCCTGACCCAAGAACAGTTTTAATGGACCA  DNA37140 3123 CTATGAGCCCAGTTACATACAGAAAAAAGGAGTGCTACCCATGTTCTCATCCTTCAGAAGA  DNA37140 3183 ATCCTGCGAACGGAGCTTCAGTAATATATCGTGGCCTTCACATGTGAGGAAGCATACTTAA  DNA37140 3243 CACTAGTTACTCTCACAATGAAGGAACCTGGACTTCACATGTGAGGAAGCACTTTAA  DNA37140 3243 CACTAGTTACTCTCACAATGAAGGACCTGGAATGAAAAAATCTGTGTCTAAACAAGTCCTC  P_AAX52264 2881 CACTAGTTACTCTCACAATGAAGGACCTGGAATGAAAAAATCTGTGTCTAAACAAGTCCTC  DNA37140 3303 TTTAGATTTTAGTGCAAATCCAGAGCCAGCGTCGGTTGCCCTCGAGTAATTTCTTTC	_AAX52264 2461 ATCTCAGGGAACGTTAGCTGACAGGCAGGATGGGTACGTGTCTTCAGAAAGTGGAAGCCA	X52264
DNA37140 2943 GACCTGCCATATTGACAATAGCAGTGAAGCTGATGTGGAAGCTGCCACAGATCTGTTCCT  P_AAX52264 2581 GACCTGCCATATTGACAATAGCAGTGAAGCTGATGTGGAAGCTGCCACAGATCTGTTCCT  DNA37140 3003 TTGTCCGTTTTTGGGATCCACAGGCCCTATGTATTTGAAGGGAAATGTGTATGGCTCAGA  P_AAX52264 2641 TTGTCCGTTTTTGGGATCCACAGGCCCTATGTATTTGAAGGGAAATGTGTATGGCTCAGA  DNA37140 3063 TCCTTTTGAAACATATCATACAGGTTGCAGTCCTGACCCAAGAACAGTTTTAATGGACCA  DNA37140 3123 CTATGAGCCCAGTTACATAAAGAAAAAGGAGTGCTACCCATGTTCTCATCCTTCAGAAGA  P_AAX52264 2761 CTATGAGCCCAGTTACATAAAGAAAAAGGAGTGCTACCCATGTTCTCATCCTTCAGAAGA  DNA37140 3183 ATCCTGCGAACGGAGCTTCAGTAATATATCGTGGCCTTCACATGTGAGGAAGCTACTTAA  P_AAX52264 2821 ATCCTGCGAACGGAGCTTCAGTAATATATCGTGGCCTTCACATGTGAGGAAGCTACTTAA  DNA37140 3243 CACTAGTTACTCTCACAATGAAGGACCTGGAATGAAAAATCTGTGTCTAAACAAGTCCTC  DNA37140 3303 TTTAGATTTTAGTGCAAATCCAGAGCCAGGGTCGGTTGCCTCGAGTAATTCTTCATGGG  DNA37140 3303 TTTAGATTTTAGTGCAAATCCAGAGCCAGGGTCGGTTGCCTCGAGTAATTCTTTCATGGG		A37140
P_AAX52264 2581 GACCTGCCATATTGACAATAGCAGTGAAGCTGATGTGGAAGCTGCCACAGATCTGTTCCT  DNA37140 3003 TTGTCCGTTTTTGGGATCCACAGGCCCTATGTATTTGAAGGGAAATGTGTATGGCTCAGA  P_AAX52264 2641 TTGTCCGTTTTTGGGATCCACAGGCCCTATGTATTTGAAGGGAAATGTGTATGGCTCAGA  DNA37140 3063 TCCTTTTGAAACATATCATACAGGTTGCAGTCCTGACCCAAGAACAGTTTTAATGGACCA  P_AAX52264 2701 TCCTTTTGAAACATATCATACAGGTTGCAGTCCTGACCCAAGAACAGTTTTAATGGACCA  DNA37140 3123 CTATGAGCCCAGTTACATAAAGAAAAAGGAGTGCTACCCATGTTCTCATCCTTCAGAAGA  DNA37140 3183 ATCCTGCGAACGGAGCTTCAGTAATATATCGTGGCCTTCACATGTGAGGAAGCTACTTAA  DNA37140 3281 ATCCTGCGAACGGAGCTTCAGTAATATATCGTGGCCTTCACATGTGAGGAAGCTACTTAA  DNA37140 3243 CACTAGTTACTCTCACAATGAAGGACCTGGAATGAAAAAATCTGTGTCTAAACAAGTCCTC  P_AAX52264 2881 CACTAGTTACTCTCACAATGAAGGACCTGGAATGAAAAAAATCTGTGTCTAAACAAGTCCTC  DNA37140 3303 TTTAGATTTTACTGCAAATCCAGAGCCAGCGTCGGTTGCCTCGAGTAATTCTTTCATGGG  DNA37140 3303 TTTAGATTTTACTGCAAATCCAGAGCCAGCGTCGGTTGCCTCGAGTAATTCTTTCATGGG  DNA37140 3303 TTTAGATTTTACTGCAAATCCAGAGCCAGCGTCGGTTGCCTCGAGTAATTCTTTCATGGG	_AAX52264 2521 CCACCAGTTTGTCACATCTTCAGGTGCTGGATTTTTCTTACCACAACATGACAGTAGTGG	X52264
P_AAX52264 2581 GACCTGCCATATTGACAATAGCAGTGAAGCTGATGTGGAAGCTGCCACAGATCTGTTCCT  DNA37140 3003 TTGTCCGTTTTTGGGATCCACAGGCCCTATGTATTTGAAGGGAAATGTGTATGGCTCAGA  P_AAX52264 2641 TTGTCCGTTTTTGGGATCCACAGGCCCTATGTATTTGAAGGGAAATGTGTATGGCTCAGA  DNA37140 3063 TCCTTTTGAAACATATCATACAGGTTGCAGTCCTGACCCAAGAACAGTTTTAATGGACCA  ********************************		A37140
P_AAX52264 2641 TTGTCCGTTTTTGGGATCCACAGGCCCTATGTATTTGAAGGAAAAATGTGTATGGCTCAGA  DNA37140 3063 TCCTTTTGAAACATATCATACAGGTTGCAGTCCTGACCCAAGAACAGTTTTAATGGACCA *********************************		X52264
DNA37140 3063 TCCTTTTGAAACATATCATACAGGTTGCAGTCCTGACCCAAGAACAGTTTTAATGGACCA  P_AAX52264 2701 TCCTTTTGAAACATATCATACAGGTTGCAGTCCTGACCCAAGAACAGTTTTAATGGACCA  DNA37140 3123 CTATGAGCCCAGTTACATAAAGAAAAAGGAGTGCTACCCATGTTCTCATCCTTCAGAAGA  DNA37140 3183 ATCCTGCGAACGGAGCTTCAGTAATATATCGTGGCCTTCACATGTGAGGAAGCTACTTAA  P_AAX52264 2821 ATCCTGCGAACGGAGCTTCAGTAATATATCGTGGCCTTCACATGTGAGGAAGCTACTTAA  DNA37140 3243 CACTAGTTACTCTCACAATGAAGGACCTGGAATGAAAAATCTGTGTCTCTAAACAAGTCCTC  AAX52264 2881 CACTAGTTACTCTCACAATGAAGGACCTGGAATGAAAAATCTGTGTCTAAACAAGTCCTC  DNA37140 3303 TTTAGATTTTAGTGCAAATCCAGAGCCAGCGTCGGTTGCCTCGAGTAATTCTTTCATGGG  TTTAGATTTTAGTGCAAATCCAGAGCCAGCTTCGGTTGCCTCGAGTAATTCTTTCATGGG  TTTAGATTTTAGTGCAAATCCAGAGCCAGCGTCGGTTGCCTCGAGTAATTCTTTCATGGG  TTTAGATTTTTAGTGCAAATCCAGAGCCAGCGTCGGTTGCCTCGAGTAATTCTTTCATGGG  TTTAGATTTTAGTGCAAATCCAGAGCCAGCGTCGGTTGCCTCGAGTAATTCTTTCATGGG  TTTAGATTTTAGTGCAAATCCAGAGCCAGCGTCGGTTGCCTCGAGTAATTCTTTCATGGG  TTTAGATTTTAGTGCAAATCCAGAGCCAGCGTCGGTTGCCTCGAGTAATTCTTTCATGGG  TTTAGATTTTAGTGCAAATCCAGAGCCAGCGTCGGTTGCCTCGAGAGCAGAGCAGAGAGAATCCAGAGCCAGCGTCGGTTGCCTCGAGTAATTCTTTCATGGG  TTTAGATTTTAGTGCAAATCCAGAGCCAGCGTCGGTTGCCTCGAGTAATTCTTTCATGGG  TTTAGATTTTAGTGCAAATCCAGAGCCAGCGTCGGTTGCCTCGAGAGCAGAGCAGAGCAGAGAGAAATCCAGAGCAGAGCTAGAATCCAGAGCAGAGCAGAGCTAGAATACAAATCCAGAGACCAGAGCAGAGAGAATCCAGAGCAGAGCAGAGCTAGAATCCAGAGCAGAGAGAATCAAATCAGAGAGAATCAAATCA		A37140
P_AAX52264 2701 TCCTTTTGAAACATATCATACAGGTTGCAGTCCTGACCCAAGAACAGTTTTAATGGACCA  DNA37140 3123 CTATGAGCCCAGTTACATAAAGAAAAAGGAGTGCTACCCATGTTCTCATCCTTCAGAAGA  *******************************	_AAX52264 2641 TTGTCCGTTTTTGGGATCCACAGGCCCTATGTATTTGAAGGGAAATGTGTATGGCTCAGA	X52264
DNA37140 3123 CTATGAGCCCAGTTACATAAAGAAAAAGGAGTGCTACCCATGTTCTCATCCTTCAGAAGA  *******************************		A37140
P_AAX52264 2761 CTATGAGCCCAGTTACATAAAGAAAAAGGAGTGCTACCCATGTTCTCATCCTTCAGAAGA  DNA37140 3183 ATCCTGCGAACGGAGCTTCAGTAATATATCGTGGCCTTCACATGTGAGGAAGCTACTTAA  ******************************	_AAX52264 2701 TCCTTTTGAAACATATCATACAGGTTGCAGTCCTGACCCAAGAACAGTTTTAATGGACCA	X52264
DNA37140 3183 ATCCTGCGAACGGAGCTTCAGTAATATATCGTGGCCTTCACATGTGAGGAAGCTACTTAA  ******************************		437140
P_AAX52264 2821 ATCCTGCGAACGGAGCTTCAGTAATATATCGTGGCCTTCACATGTGAGGAAGCTACTTAA  DNA37140 3243 CACTAGTTACTCTCACAATGAAGGACCTGGAATGAAAAATCTGTGTCTAAACAAGTCCTC *********************************	_AAX52264 2761 CTATGAGCCCAGTTACATAAAGAAAAAGGAGTGCTACCCATGTTCTCATCCTTCAGAAGA	X52264
DNA37140 3243 CACTAGTTACTCTCACAATGAAGGACCTGGAATGAAAAATCTGTGTCTAAACAAGTCCTC  ********************************		437140
P_AAX52264 2881 CACTAGTTACTCTCACAATGAAGGACCTGGAATGAAAAATCTGTGTCTAAACAAGTCCTC  DNA37140 3303 TTTAGATTTTAGTGCAAATCCAGAGCCAGCGTCGGTTGCCTCGAGTAATTCTTTCATGGG		X52264
DNA37140 3303 TTTAGATTTTAGTGCAAATCCAGAGCCAGCGTCGGTTGCCTCGAGTAATTCTTTCATGGG		437140
************		X52264
		437140
		X52264

```
3363 TACCTTTGGAAAAGCTCTCAGGAGACCTCACCTAGATGCCTATTCAAGCTTTGGACAGCC
 DNA37140
             ***********
         3001 TACCTTTGGAAAAGCTCTCAGGAGACCTCACCTAGATGCCTATTCAAGCTTTGGACAGCC
P AAX52264
         3423 ATCAGATTGTCAGCCAAGAGCCTTTTATTTGAAAGCTCATTCTTCCCCAGACTTGGACTC
 DNA37140
            P AAX52264
         3061 ATCAGATTGTCAGCCAAGAGCCTTTTATTTGAAAGCTCATTCTTCCCCAGACTTGGACTC
         3483 TGGGTCAGAGGAAGATGGGAAAGAAAGGACAGATTTTCAGGAAGAAAATCACATTTGTAC
 DNA37140
            P AAX52264
         3121 TGGGTCAGAGGAAGATGGGAAAGAAGGACAGATTTTCAGGAAGAAAATCACATTTGTAC
 DNA37140
         3543 CTTTAAACAGACTTTAGAAAACTACAGGACTCCAAATTTTCAGTCTTATGACTTGGACAC
            P AAX52264
         3181 CTTTAAACAGACTTTAGAAAACTACAGGACTCCAAATTTTCAGTCTTATGACTTGGACAC
 DNA37140
         P AAX52264
 DNA37140
         3663 AAAGAGAGAATCTTATGTTTTTTAAATGGAGTTATGAATTTTAAAAGGATAAAAATGC
         3301 AAAGAGAGAATCTTATGTTTTTTAAATGGAGTTATGAATTTTAAAAGGATAAAAATGC
P AAX52264
         3723 TTTATTTATACAGATGAACCAAAATTACAAAAAGTTATGAAAATTTTTATACTGGGAATG
 DNA37140
            P AAX52264
         3361 TTTATTTATACAGATGAACCAAAATTACAAAAAGTTATGAAAATTTTTATACTGGGAATG
 DNA37140
         3783 ATGCTCATATAAGAATACCTTTTTAAACTATTTTTTAACTTTGTTTTATGCAAAAAAGTA
P AAX52264
         3421 ATGCTCATATAAGAATACCTTTTTAAACTATTTTTTAACTTTGTTTTATGCAAAAAAGTA
        3843 TCTTACGTAAATTAATGATATAAATCATGATTATTTTATGTATTTTATAATGCCAGATT
 DNA37140
            P AAX52264
        3481 TCTTACGTAAATTAATGATATAAATCATGATTATTTTATGTATTTTTATAATGCCAGATT
        3903 TCTTTTTATGGAAAATGAGTTACTAAAGCATTTTAAATAATACCTGCCTTGTACCATTTT
 DNA37140
            ***********
        3541 TCTTTTTATGGAAAATGAGTTACTAAAGCATTTTAAATAATACCTGCCTTGTACCATTTT
P AAX52264
 DNA37140
        3963 TTAAATAGAAGTTACTTCATTATATTTTGCACATTATATTTAATAAAATGTGTCAATTTG
            3601 TTAAATAGAAGTTACTTCATTATATTTTGCACATTATATTTAATAAAATGTGTCAATTTG
P AAX52264
 DNA37140
        4023 AA
P AAX52264
        3661 AA
>11 P AAX37725 Human PRO335 DNA fragment #2. (3598 bp) [1 seg]
Score = 3542 (7022 bits), Expect = 0.0
Identities = 3542/3542 (100%), at 483,57-4024,3598, Strand +/+
         483 GGACTTAAGTCACAACAGATTATCTTTCATCAAGGCAAGTTCCATGAGCCACCTTCAAAG
 DNA37140
            P AAX37725
          57 GGACTTAAGTCACAACAGATTATCTTTCATCAAGGCAAGTTCCATGAGCCACCTTCAAAG
         543 CCTTCGAGAAGTGAAACTGAACAACAATGAATTGGAGACCATTCCAAATCTGGGACCAGT
 DNA 37140
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P_AAX37725	117	CCTTCGAGAAGTGAAACTGAACAACAATGAATTGGAGACCATTCCAAATCTGGGACCAGT
DNA37140	603	CTCGGCAAATATTACACTTCTCTCCTTGGCTGGAAACAGGATTGTTGAAATACTCCCTGA
P_AAX37725	177	CTCGGCAAATATTACACTTCTCCTTGGCTGGAAACAGGATTGTTGAAATACTCCCTGA
DNA37140	663	ACATCTGAAAGAGTTTCAGTCCCTTGAAACTTTGGACCTTAGCAGCAACAATATTTCAGA
P_AAX37725	237	ACATCTGAAAGAGTTTCAGTCCCTTGAAACTTTGGACCTTAGCAGCAACAATATTTCAGA
DNA37140	723	GCTCCAAACTGCATTTCCAGCCCTACAGCTCAAATATCTGTATCTCAACAGCAACCGAGT
P_AAX37725	297	GCTCCAAACTGCATTTCCAGCCCTACAGCTCAAATATCTGTATCTCAACAGCAACCGAGT
DNA37140	783	CACATCAATGGAACCTGGGTATTTTGACAATTTGGCCAACACACTCCTTGTGTTAAAGCT
P_AAX37725	357	
DNA37140	843	GAACAGGAACCGAATCTCAGCTATCCCACCCAAGATGTTTAAACTGCCCCAACTGCAACA
P_AAX37725	417	GAACAGGAACCGAATCTCAGCTATCCCACCCAAGATGTTTAAACTGCCCCAACTGCAACA
DNA37140	903	TCTCGAATTGAACCGAAACAAGATTAAAAATGTAGATGGACTGACATTCCAAGGCCTTGG
P_AAX37725	477	TCTCGAATTGAACCGAAACAAGATTAAAAATGTAGATGGACTGACATTCCAAGGCCTTGG
DNA37140	963	TGCTCTGAAGTCTCTGAAAATGCAAAGAAATGGAGTAACGAAACTTATGGATGG
P_AAX37725	537	TGCTCTGAAGTCTCTGAAAATGCAAAGAAATGGAGTAACGAAACTTATGGATGG
DNA37140	1023	TTGGGGGCTGAGCAACATGGAAATTTTGCAGCTGGACCATAACAACCTAACAGAGATTAC
P_AAX37725	597	
DNA37140	1083	CAAAGGCTGGCTTTACGGCTTGCTGATGCTGCAGGAACTTCATCTCAGCCAAAATGCCAT
P_AAX37725	657	${\tt CAAAGGCTGGCTTTACGGCTTGCTGATGCTGCAGGAACTTCATCTCAGCCAAAATGCCATCATCTCAGCCAAAATGCCATCATCTCAGCCAAAATGCCATCATCTCAGCCAAAATGCCATCATCTCAGCCAAAATGCCATCATCTCAGCCAAAATGCCATCATCTCAGCCAAAATGCCATCATCATCTCAGCCAAAAATGCCATCATCATCTCAGCCAAAAATGCCATCATCATCAGCCAAAAATGCCATCATCAGCCAAAAATGCCATCATCAGCAAAAATGCCATCATCAGCAAAAATGCCATCATCAGCAAAAATGCCATCAGCAAAAATGCCATCAGCAAAAATGCCATCAGCAAAAATGCCATCAGCAAAAAATGCCATCAGCAAAAATGCCATCAGCAAAAATGCCATCAGCAAAAATGCCATCAGCAAAAATGCCATCAGCAAAAATGCCATCAGCAAAAATGCCATCAGCAAAAATGCCATCAGCAAAAATGCCATCAGCAAAAATGCCATCAGCAAAAATGCCATCAGCAAAAATGCCATCAGCAAAAATGCCATCAGCAAAAAATGCCATCAGCAAAAAATGCCATCAGCAAAAAATGCCATCAGCAAAAAATGCCATCAGCAAAAAATGCCATCAGCAAAAAATGCCATCAGCAAAAAATGCCATCAGCAAAAAATGCCATCAGCAAAAAATGCCATCAGCAAAAAATGCCATCAGCAAAAAATGCCATCAGCAAAAAATGCCAATCAGCAAAAAATGCCATCAGCAAAAAAATGCCAATCAGCAAAAAAAA$
DNA37140	1143	CAACAGGATCAGCCCTGATGCCTGGGAGTTCTGCCAGAAGCTCAGTGAGCTGGACCTAAC
P_AAX37725	717	CAACAGGATCAGCCCTGATGCCTGGGAGTTCTGCCAGAAGCTCAGTGAGCTGGACCTAAC
DNA37140	1203	TTTCAATCACTTATCAAGGTTAGATGATTCAAGCTTCCTTGGCCTAAGCTTACTAAATAC
P_AAX37725	777	TTTCAATCACTTATCAAGGTTAGATGATTCAAGCTTCCTTGGCCTAAGCTTACTAAATAC
DNA37140	1263	ACTGCACATTGGGAACAACAGAGTCAGCTACATTGCTGATTGTGCCTTCCGGGGGCTTTC
P_AAX37725	837	${\tt ACTGCACATTGGGAACAACAGAGTCAGCTACATTGCTGATTGTGCCTTCCGGGGGCTTTCCGGGGGGCTTTCCGGGGGG$
DNA37140	1323	CAGTTTAAAGACTTTGGATCTGAAGAACAATGAAATTTCCTGGACTATTGAAGACATGAA
P_AAX37725	897	CAGTTTAAAGACTTTGGATCTGAAGAACAATGAAATTTCCTGGACTATTGAAGACATGAA
DNA37140	1383	TGGTGCTTTCTCTGGGCTTGACAAACTGAGGCGACTGATACTCCAAGGAAATCGGATCCG

P_AAX37725	957	TGGTGCTTTCTCTGGGCTTGACAAACTGAGGCGACTGATACTCCAAGGAAATCGGATCCG
DNA37140		TTCTATTACTAAAAAAGCCTTCACTGGTTTGGATGCATTGGAGCATCTAGACCTGAGTGA
P_AAX37725	1017	TTCTATTACTAAAAAAGCCTTCACTGGTTTGGATGCATTGGAGCATCTAGACCTGAGTGA
DNA37140	1503	CAACGCAATCATGTCTTTACAAGGCAATGCATTTTCACAAATGAAGAAACTGCAACAATT
P_AAX37725	1077	CAACGCAATCATGTCTTTACAAGGCAATGCATTTTCACAAATGAAGAAACTGCAACAATT
DNA37140	1563	GCATTTAAATACATCAAGCCTTTTGTGCGATTGCCAGCTAAAATGGCTCCCACAGTGGGT ********************************
P_AAX37725	1137	GCATTTAAATACATCAAGCCTTTTGTGCGATTGCCAGCTAAAATGGCTCCCACAGTGGGT
DNA37140	1623	GGCGGAAAACAACTTTCAGAGCTTTGTAAATGCCAGTTGTGCCCATCCTCAGCTGCTAAA **********************************
P_AAX37725	1197	GGCGGAAAACAACTTTCAGAGCTTTGTAAATGCCAGTTGTGCCCATCCTCAGCTGCTAAA
DNA37140	1683	AGGAAGAAGCATTTTTGCTGTTAGCCCAGATGGCTTTGTGTGTG
P_AAX37725	1257	AGGAAGAAGCATTTTTGCTGTTAGCCCAGATGGCTTTGTGTGTG
DNA37140	1743	CCAGATCACGGTTCAGCCAGAAACACAGTCGGCAATAAAAGGTTCCAATTTGAGTTTCAT
P_AAX37725	1317	CCAGATCACGGTTCAGCCAGAAACACAGTCGGCAATAAAAGGTTCCAATTTGAGTTTCAT
DNA37140	1803	CTGCTCAGCTGCCAGCAGCAGTGATTCCCCAATGACTTTTGCTTGGAAAAAAAGACAATGA
P_AAX37725	1377	CTGCTCAGCTGCCAGCAGCAGTGATTCCCCCAATGACTTTTGCTTGGAAAAAAAGACAATGA
DNA37140	1863	ACTACTGCATGATGCTGAAATGGAAAATTATGCACACCTCCGGGCCCAAGGTGGCGAGGT **********************************
P_AAX37725	1437	ACTACTGCATGATGCTGAAATGGAAAATTATGCACACCTCCGGGCCCAAGGTGGCGAGGT
DNA37140	1923	GATGGAGTATACCACCATCCTTCGGCTGCGCGAGGTGGAATTTGCCAGTGAGGGGAAATA *************************
P_AAX37725	1497	GATGGAGTATACCACCATCCTTCGGCTGCGCGAGGTGGAATTTGCCAGTGAGGGGAAATA
DNA37140	1983	TCAGTGTGTCATCTCCAATCACTTTGGTTCATCCTACTCTGTCAAAGCCAAGCTTACAGT
P_AAX37725	1557	TCAGTGTGTCATCTCCAATCACTTTGGTTCATCCTACTCTGTCAAAGCCAAGCTTACAGT
DNA37140	2043	AAATATGCTTCCCTCATTCACCAAGACCCCCATGGATCTCACCATCCGAGCTGGGGCCAT ***********************************
P_AAX37725	1617	AAATATGCTTCCCTCATTCACCAAGACCCCCATGGATCTCACCATCCGAGCTGGGGCCAT
DNA37140	2103	GGCACGCTTGGAGTGTGCTGTGGGGCACCCAGCCCCCAGATAGCCTGGCAGAAGGA *****************************
P_AAX37725	1677	GGCACGCTTGGAGTGTGCTGTGGGGGCACCCAGCCCCCAGATAGCCTGGCAGAAGGA
DNA37140	2163	TGGGGGCACAGACTTCCCAGCTGCACGGGAGAGACGCATGCAT
P_AAX37725	1737	TGGGGGCACAGACTTCCCAGCTGCACGGGAGAGACGCATGCAT
DNA37140	2223	CGTGTTCTTTATCGTGGATGTGAAGATAGAGGACATTGGGGTATACAGCTGCACAGCTCA
P_AAX37725	1797	CGTGTTCTTTATCGTGGATGTGAAGATAGAGGACATTGGGGTATACAGCTGCACAGCTCA

DNA37140	2283	GAACAGTGCAGGAAGTATTTCAGCAAATGCAACTCTGACTGTCCTAGAAACACCATCATT
P_AAX37725	1857	
DNA37140	2343	TTTGCGGCCACTGTTGGACCGAACTGTAACCAAGGGAGAAACAGCCGTCCTACAGTGCAT
P_AAX37725	1917	TTTGCGGCCACTGTTGGACCGAACTGTAACCAAGGGAGAAACAGCCGTCCTACAGTGCAT
DNA37140	2403	TGCTGGAGGAAGCCCTCCCCCTAAACTGAACTGGACCAAAGATGATAGCCCATTGGTGGT
P_AAX37725	1977	TGCTGGAGGAAGCCCTCCCCCTAAACTGAACTGGACCAAAGATGATAGCCCATTGGTGGT
DNA37140	2463	AACCGAGAGGCACTTTTTTGCAGCAGGCAATCAGCTTCTGATTATTGTGGACTCAGATGT *********************************
P_AAX37725	2037	AACCGAGAGGCACTTTTTTGCAGCAGGCAATCAGCTTCTGATTATTGTGGACTCAGATGT
DNA37140	2523	CAGTGATGCTGGGAAATACACATGTGAGATGTCTAACACCCTTGGCACTGAGAGAGA
P_AAX37725	2097	CAGTGATGCTGGGAAATACACATGTGAGATGTCTAACACCCTTGGCACTGAGAGAGA
DNA37140	2583	CGTGCGCCTCAGTGTGATCCCCACTCCAACCTGCGACTCCCCTCAGATGACAGCCCCATC
P_AAX37725	2157	CGTGCGCCTCAGTGTGATCCCCACTCCAACCTGCGACTCCCCTCAGATGACAGCCCCATC
DNA37140	2643	GTTAGACGATGACGGATGGGCCACTGTGGGTGTCGTGATCATAGCCGTGGTTTGCTGTGT **********************
P_AAX37725	2217	GTTAGACGATGACGGATGGGCCACTGTGGGTGTCGTGATCATAGCCGTGGTTTGCTGTGT
DNA37140	2703	GGTGGCACGTCACTCGTGTGGGTGGTCATCATATACCACACAGGCGGAGGAATGAAGA **************************
P_AAX37725	2277	GGTGGCACGTCACTCGTGTGGGTGGTCATCATATACCACACAAGGCGGAGGAATGAAGA
DNA37140	2763	TTGCAGCATTACCAACACAGATGAGACCAACTTGCCAGCAGATATTCCTAGTTATTTGTC *********************************
P_AAX37725	2337	TTGCAGCATTACCAACACAGATGAGACCAACTTGCCAGCAGATATTCCTAGTTATTTGTC
DNA37140	2823	ATCTCAGGGAACGTTAGCTGACAGGCAGGATGGGTACGTGTCTTCAGAAAGTGGAAGCCA *******************************
P_AAX37725	2397	ATCTCAGGGAACGTTAGCTGACAGGCAGGATGGGTACGTGTCTTCAGAAAGTGGAAGCCA
DNA37140	2883	CCACCAGTTTGTCACATCTTCAGGTGCTGGATTTTTCTTACCACAACATGACAGTAGTGG *******************************
P_AAX37725	2457	CCACCAGTTTGTCACATCTTCAGGTGCTGGATTTTTCTTACCACAACATGACAGTAGTGG
DNA37140	2943	GACCTGCCATATTGACAATAGCAGTGAAGCTGATGTGGAAGCTGCCACAGATCTGTTCCT ******************************
P_AAX37725	2517	GACCTGCCATATTGACAATAGCAGTGAAGCTGATGTGGAAGCTGCCACAGATCTGTTCCT
DNA37140		TTGTCCGTTTTTGGGATCCACAGGCCCTATGTATTTGAAGGGAAATGTGTATGGCTCAGA ***********************************
P_AAX37725	2577	$\tt TTGTCCGTTTTTGGGGATCCACAGGCCCTATGTATTTGAAGGGAAATGTGTATTGGCTCAGA$
DNA37140	3063	TCCTTTTGAAACATATCATACAGGTTGCAGTCCTGACCCAAGAACAGTTTTAATGGACCA *********************************
P_AAX37725	2637	TCCTTTTGAAACATATCATACAGGTTGCAGTCCTGACCCAAGAACAGTTTTAATGGACCA

DNA37140	3123	CTATGAGCCCAGTTACATAAAGAAAAAGGAGTGCTACCCATGTTCTCATCCTTCAGAAGA
P_AAX37725	2697	CTATGAGCCCAGTTACATAAAGAAAAAGGAGTGCTACCCATGTTCTCATCCTTCAGAAGA
DNA37140	3183	ATCCTGCGAACGGAGCTTCAGTAATATATCGTGGCCTTCACATGTGAGGAAGCTACTTAA
P_AAX37725	2757	ATCCTGCGAACGGAGCTTCAGTAATATATCGTGGCCTTCACATGTGAGGAAGCTACTTAA
DNA37140	3243	CACTAGTTACTCTCACAATGAAGGACCTGGAATGAAAAATCTGTGTCTAAACAAGTCCTC
P_AAX37725	2817	CACTAGTTACTCTCACAATGAAGGACCTGGAATGAAAAATCTGTGTCTAAACAAGTCCTC
DNA37140	3303	TTTAGATTTTAGTGCAAATCCAGAGCCAGCGTCGGTTGCCTCGAGTAATTCTTTCATGGG **********************************
P_AAX37725	2877	TTTAGATTTTAGTGCAAATCCAGAGCCAGCGTCGGTTGCCTCGAGTAATTCTTTCATGGG
DNA37140	3363	TACCTTTGGAAAAGCTCTCAGGAGACCTCACCTAGATGCCTATTCAAGCTTTGGACAGCC
P_AAX37725	2937	TACCTTTGGAAAAGCTCTCAGGAGACCTCACCTAGATGCCTATTCAAGCTTTGGACAGCC
DNA37140	3423	ATCAGATTGTCAGCCAAGAGCCTTTTATTTGAAAGCTCATTCTTCCCCAGACTTGGACTC
P_AAX37725	2997	ATCAGATTGTCAGCCAAGAGCCTTTTATTTGAAAGCTCATTCTTCCCCAGACTTGGACTC
DNA37140	3483	TGGGTCAGAGGAAGAAGGAAGAAGGACAGATTTTCAGGAAGAAAATCACATTTGTAC
P_AAX37725	3057	TGGGTCAGAGGAAGAAGGAAAGGACAGATTTTCAGGAAGAAAATCACATTTGTAC
DNA37140	3543	CTTTAAACAGACTTTAGAAAACTACAGGACTCCAAATTTTCAGTCTTATGACTTGGACAC
P_AAX37725	3117	CTTTAAACAGACTTTAGAAAACTACAGGACTCCAAATTTTCAGTCTTATGACTTGGACAC
DNA37140	3603	ATAGACTGAATGAGACCAAAGGAAAAGCTTAACATACTACCTCAAGTGAACTTTTATTTA
P_AAX37725	3177	ATAGACTGAATGAGACCAAAGGAAAAGCTTAACATACTACCTCAAGTGAACTTTTATTTA
DNA37140	3663	AAAGAGAGAGAATCTTATGTTTTTTAAATGGAGTTATGAATTTTAAAAGGATAAAAATGC
P_AAX37725	3237	AAAGAGAGAGAATCTTATGTTTTTTAAATGGAGTTATGAATTTTAAAAGGATAAAAATGC
DNA37140	3723	TTTATTTATACAGATGAACCAAAATTACAAAAAGTTATGAAAATTTTTATACTGGGAATG
P_AAX37725	3297	TTTATTTATACAGATGAACCAAAATTACAAAAAGTTATGAAAATTTTTATACTGGGAATG
DNA37140	3783	ATGCTCATATAAGAATACCTTTTTAAACTATTTTTTAACTTTGTTTTATGCAAAAAGTA
P_AAX37725	3357	ATGCTCATATAAGAATACCTTTTTAAACTATTTTTTAACTTTTTTTATGCAAAAAAGTA
DNA37140	3843	TCTTACGTAAATTAATGATATAAATCATGATTATTTTATGTATTTTTATAATGCCAGATT **********************************
P_AAX37725	3417	TCTTACGTAAATTAATGATATAAATCATGATTATTTTATGTATTTTTATAATGCCAGATT
DNA37140	3903	TCTTTTTATGGAAAATGAGTTACTAAAGCATTTTAAATAATACCTGCCTTGTACCATTTT ******************************
P_AAX37725	3477	TCTTTTTATGGAAAATGAGTTACTAAAGCATTTTAAATAATACCTGCCTTGTACCATTTT
DNA37140	3963	TTAAATAGAAGTTACTTCATTATTTTTGCACATTATATTTAATAAAATGTGTCAATTTG

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        3537 TTAAATAGAAGTTACTTCATTATATTTTGCACATTATATTTAATAAAATGTGTCAATTTG
 DNA37140
        4023 AA
P AAX37725 3597 AA
>12 AX098383 Sequence 9 from Patent WO0119991. (3662 bp) [1 seq]
Score = 3542 (7022 bits), Expect = 0.0
Identities = 3542/3542 (100%), at 483,121-4024,3662, Strand +/+
 DNA37140
         483 GGACTTAAGTCACAACAGATTATCTTTCATCAAGGCAAGTTCCATGAGCCACCTTCAAAG
            121 GGACTTAAGTCACAACAGATTATCTTTCATCAAGGCAAGTTCCATGAGCCACCTTCAAAG
 AX098383
         543 CCTTCGAGAAGTGAAACTGAACAATGAATTGGAGACCATTCCAAATCTGGGACCAGT
 DNA37140
            181 CCTTCGAGAAGTGAAACTGAACAACAATGAATTGGAGACCATTCCAAATCTGGGACCAGT
 AX098383
         603 CTCGGCAAATATTACACTTCTCCTTGGCTGGAAACAGGATTGTTGAAATACTCCCTGA
 DNA37140
            ************
         241 CTCGGCAAATATTACACTTCTCTCCTTGGCTGGAAACAGGATTGTTGAAATACTCCCTGA
 AX098383
         663 ACATCTGAAAGAGTTTCAGTCCCTTGAAACTTTGGACCTTAGCAGCAACAATATTTCAGA
 DNA37140
            ************
         301 ACATCTGAAAGAGTTTCAGTCCCTTGAAACTTTGGACCTTAGCAGCAACAATATTTCAGA
 AX098383
         723 GCTCCAAACTGCATTTCCAGCCCTACAGCTCAAATATCTGTATCTCAACAGCAACCGAGT
 DNA37140
            ***********
         361 GCTCCAAACTGCATTTCCAGCCCTACAGCTCAAATATCTGTATCTCAACAGCAACCGAGT
 AX098383
         783 CACATCAATGGAACCTGGGTATTTTGACAATTTGGCCAACACACCTCCTTGTGTTAAAGCT
 DNA37140
            421 CACATCAATGGAACCTGGGTATTTTGACAATTTGGCCAACACACTCCTTGTGTTAAAGCT
 AX098383
         843 GAACAGGAACCGAATCTCAGCTATCCCACCCAAGATGTTTAAACTGCCCCAACTGCAACA
 DNA37140
            481 GAACAGGAACCGAATCTCAGCTATCCCACCCAAGATGTTTAAACTGCCCCAACTGCAACA
 AX098383
         903 TCTCGAATTGAACCGAAACAAGATTAAAAATGTAGATGGACTGACATTCCAAGGCCTTGG
 DNA37140
            ***********
         541 TCTCGAATTGAACCGAAACAAGATTAAAAATGTAGATGGACTGACATTCCAAGGCCTTGG
 AX098383
         DNA37140
            ***********
         AX098383
        1023 TTGGGGGCTGAGCAACATGGAAATTTTGCAGCTGGACCATAACAACCTAACAGAGATTAC
 DNA37140
            **********
 AX098383
         661 TTGGGGGCTGAGCAACATGGAAATTTTGCAGCTGGACCATAACAACCTAACAGAGATTAC
        1083 CAAAGGCTGGCTTTACGGCTTGCTGATGCTGCAGGAACTTCATCTCAGCCAAAATGCCAT
 DNA 37140
            **********
         721 CAAAGGCTGGCTTTACGGCTTGCTGATGCTGCAGGAACTTCATCTCAGCCAAAATGCCAT
 AX098383
        1143 CAACAGGATCAGCCCTGATGCCTGGGAGTTCTGCCAGAAGCTCAGTGAGCTGGACCTAAC
 DNA37140
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AXU98383	/81	. CAACAGGATCAGCCCTGATGCCTGGGAGTTCTGCCAGAAGCTCAGTGAGCTGGACCTAAC
DNA37140	1203	TTTCAATCACTTATCAAGGTTAGATGATTCAAGCTTCCTTGGCCTAAGCTTACTAAATAG
AX098383	841	TTTCAATCACTTATCAAGGTTAGATGATTCAAGCTTCCTTGGCCTAAGCTTACTAAATAC
DNA37140	1263	ACTGCACATTGGGAACAACAGAGTCAGCTACATTGCTGATTGTGCCTTCCGGGGGCTTTC
AX098383	901	ACTGCACATTGGGAACAACAGAGTCAGCTACATTGCTGATTGTGCCTTCCGGGGGCTTTC
DNA37140	1323	CAGTTTAAAGACTTTGGATCTGAAGAACAATGAAATTTCCTGGACTATTGAAGACATGAA
AX098383	961	CAGTTTAAAGACTTTGGATCTGAAGAACAATGAAATTTCCTGGACTATTGAAGACATGAA
DNA37140	1383	TGGTGCTTTCTCTGGGCTTGACAAACTGAGGCGACTGATACTCCAAGGAAATCGGATCCG
AX098383	1021	TGGTGCTTTCTCTGGGCTTGACAAACTGAGGCGACTGATACTCCAAGGAAATCGGATCCG
DNA37140	1443	TTCTATTACTAAAAAAGCCTTCACTGGTTTGGATGCATTGGAGCATCTAGACCTGAGTGA
AX098383	1081	TTCTATTACTAAAAAAGCCTTCACTGGTTTGGATGCATTGGAGCATCTAGACCTGAGTGA
DNA37140	1503	CAACGCAATCATGTCTTTACAAGGCAATGCATTTTCACAAATGAAGAAACTGCAACAATT
AX098383	1141	CAACGCAATCATGTCTTTACAAGGCAATGCATTTTCACAAATGAAGAAACTGCAACAATT
DNA37140	1563	GCATTTAAATACATCAAGCCTTTTGTGCGATTGCCAGCTAAAATGGCTCCCACAGTGGGT
AX098383	1201	GCATTTAAATACATCAAGCCTTTTGTGCGATTGCCAGCTAAAATGGCTCCCACAGTGGGT
DNA37140	1623	GGCGGAAAACAACTTTCAGAGCTTTGTAAATGCCAGTTGTGCCCATCCTCAGCTGCTAAA
AX098383	1261	GGCGGAAAACAACTTTCAGAGCTTTGTAAATGCCAGTTGTGCCCATCCTCAGCTGCTAAA
DNA37140	1683	AGGAAGAAGCATTTTTGCTGTTAGCCCAGATGGCTTTGTGTGTG
AX098383	1321	AGGAAGAAGCATTTTTGCTGTTAGCCCAGATGGCTTTGTGTGTG
DNA37140	1743	CCAGATCACGGTTCAGCCAGAAACACAGTCGGCAATAAAAGGTTCCAATTTGAGTTTCAT
AX098383	1381	${\tt CCAGATCACGGTTCAGCCAGAAACACAGTCGGCAATAAAAGGTTCCAATTTGAGTTTCATCAGGTCAGAAAAAAAA$
DNA37140	1803	CTGCTCAGCTGCCAGCAGCAGTGATTCCCCAATGACTTTTGCTTGGAAAAAAAGACAATGA
AX098383	1441	CTGCTCAGCTGCCAGCAGCAGTGATTCCCCCAATGACTTTTGCTTGGAAAAAAAGACAATGA
DNA37140	1863	ACTACTGCATGATGCTGAAATGGAAAATTATGCACACCTCCGGGCCCAAGGTGGCGAGGT
AX098383	1501	ACTACTGCATGATGCTGAAATGGAAAATTATGCACACCTCCGGGCCCAAGGTGGCGAGGT
DNA37140	1923	GATGGAGTATACCACCATCCTTCGGCTGCGCGAGGTGGAATTTGCCAGTGAGGGGAAATA
AX098383	1561	GATGGAGTATACCACCATCCTTCGGCTGCGCGAGGTGGAATTTGCCAGTGAGGGGAAATA
DNA37140	1983	TCAGTGTGTCATCTCCAATCACTTTGGTTCATCCTACTCTGTCAAAGCCAAGCTTACAGT
AX098383	1621	TCAGTGTGTCATCTCCAATCACTTTGGTTCATCCTACTCTGTCAAAGCCAAGCTTACAGT

DNA37140	2043	AAATATGCTTCCCTCATTCACCAAGACCCCCATGGATCTCACCATCCGAGCTGGGGCCAT ***********************************
AX098383	1681	AAATATGCTTCCCTCATTCACCAAGACCCCCATGGATCTCACCATCCGAGCTGGGGCCAT
DNA37140	2103	GGCACGCTTGGAGTGTGCTGTGGGGCACCCAGCCCCCAGATAGCCTGGCAGAAGGA
AX098383	1741	GGCACGCTTGGAGTGTGCTGTGGGGCACCCAGCCCCCAGATAGCCTGGCAGAAGGA
DNA37140	2163	TGGGGGCACAGACTTCCCAGCTGCACGGGAGAGCGCATGCAT
AX098383	1801	TGGGGGCACAGACTTCCCAGCTGCACGGGAGAGACGCATGCAT
DNA37140	2223	CGTGTTCTTTATCGTGGATGTGAAGATAGAGGACATTGGGGTATACAGCTGCACAGCTCA
AX098383	1861	CGTGTTCTTTATCGTGGATGTGAAGATAGAGGACATTGGGGTATACAGCTGCACAGCTCA
DNA37140	2283	GAACAGTGCAGGAAGTATTTCAGCAAATGCAACTCTGACTGTCCTAGAAACACCATCATT
AX098383	1921	GAACAGTGCAGGAAGTATTTCAGCAAATGCAACTCTGACTGTCCTAGAAACACCATCATT
DNA37140	2343	TTTGCGGCCACTGTTGGACCGAACTGTAACCAAGGGAGAAACAGCCGTCCTACAGTGCAT
AX098383	1981	TTTGCGGCCACTGTTGGACCGAACTGTAACCAAGGGAGAAACAGCCGTCCTACAGTGCAT
DNA37140	2403	TGCTGGAGGAAGCCCTCCCCCTAAACTGAACTGGACCAAAGATGATAGCCCATTGGTGGT
AX098383	2041	TGCTGGAGGAAGCCCTCCCCCTAAACTGAACTGGACCAAAGATGATAGCCCATTGGTGGT
DNA37140	2463	AACCGAGAGGCACTTTTTTGCAGCAGGCAATCAGCTTCTGATTATTGTGGACTCAGATGT
AX098383	2101	AACCGAGAGGCACTTTTTTGCAGCAGCCAATCAGCTTCTGATTATTGTGGACTCAGATGT
DNA37140	2523	CAGTGATGCTGGGAAATACACATGTGAGATGTCTAACACCCTTGGCACTGAGAGAGGAAA
AX098383	2161	CAGTGATGCTGGGAAATACACATGTGAGATGTCTAACACCCTTGGCACTGAGAGAGA
DNA37140	2583	CGTGCGCCTCAGTGTGATCCCCACTCCAACCTGCGACTCCCCTCAGATGACAGCCCCATC
AX098383	2221	CGTGCGCCTCAGTGTGATCCCCACTCCAACCTGCGACTCCCCTCAGATGACAGCCCCATC
DNA37140	2643	GTTAGACGATGACGGATGGGCCACTGTGGGTGTCGTGATCATAGCCGTGGTTTGCTGTGT
AX098383	2281	GTTAGACGATGACGGATGGGCCACTGTGGGTGTCGTGATCATAGCCGTGGTTTGCTGTGT
DNA37140	2703	GGTGGGCACGTCACTCGTGTGGGTGGTCATCATATACCACACAAGGCGGAGGAATGAAGA
AX098383	2341	GGTGGGCACGTCACTCGTGTGGGTGGTCATCATATACCACACAAGGCGGAGGAATGAAGA
DNA37140	2763	TTGCAGCATTACCAACACAGATGAGACCAACTTGCCAGCAGATATTCCTAGTTATTTGTC
AX098383	2401	TTGCAGCATTACCAACACAGATGAGACCAACTTGCCAGCAGATATTCCTAGTTATTTGTC
DNA37140	2823	ATCTCAGGGAACGTTAGCTGACAGGCAGGATGGGTACGTGTCTTCAGAAAGTGGAAGCCA
AX098383	2461	ATCTCAGGGAACGTTAGCTGACAGGCAGGATGGGTACGTGTCTTCAGAAAGTGGAAGCCA

DNA37140	2883	CCACCAGTTTGTCACATCTTCAGGTGCTGGATTTTTCTTACCACAACATGACAGTAGTGG *******************************
AX098383	2521	CCACCAGTTTGTCACATCTTCAGGTGCTGGATTTTTCTTACCACAACATGACAGTAGTGG
DNA37140	2943	GACCTGCCATATTGACAATAGCAGTGAAGCTGATGTGGAAGCTGCCACAGATCTGTTCCT
AX098383	2581	GACCTGCCATATTGACAATAGCAGTGAAGCTGATGTGGAAGCTGCCACAGATCTGTTCCT
DNA37140	3003	TTGTCCGTTTTTGGGATCCACAGGCCCTATGTATTTGAAGGGAAATGTGTATGGCTCAGA
AX098383	2641	TTGTCCGTTTTTGGGATCCACAGGCCCTATGTATTTGAAGGGAAATGTGTATGGCTCAGA
DNA37140	3063	TCCTTTTGAAACATATCATACAGGTTGCAGTCCTGACCCAAGAACAGTTTTAATGGACCA *********************************
AX098383	2701	TCCTTTTGAAACATATCATACAGGTTGCAGTCCTGACCCAAGAACAGTTTTAATGGACCA
DNA37140	3123	CTATGAGCCCAGTTACATAAAGAAAAAGGAGTGCTACCCATGTTCTCATCCTTCAGAAGA
AX098383	2761	CTATGAGCCCAGTTACATAAAGAAAAAGGAGTGCTACCCATGTTCTCATCCTTCAGAAGA
DNA37140	3183	ATCCTGCGAACGGAGCTTCAGTAATATCGTGGCCTTCACATGTGAGGAAGCTACTTAA
AX098383	2821	$\tt ATCCTGCGAACGGAGCTTCAGTAATATCGTGGCCTTCACATGTGAGGAAGCTACTTAA$
DNA37140	3243	CACTAGTTACTCTCACAATGAAGGACCTGGAATGAAAAATCTGTGTCTAAACAAGTCCTC
AX098383	2881	CACTAGTTACTCTCACAATGAAGGACCTGGAATGAAAAATCTGTGTCTAAACAAGTCCTC
DNA37140	3303	TTTAGATTTTAGTGCAAATCCAGAGCCAGCGTCGGTTGCCTCGAGTAATTCTTTCATGGG **********************************
AX098383	2941	${\tt TTTAGATTTTAGTGCAAATCCAGAGCCAGCGTCGGTTGCCTCGAGTAATTCTTTCATGGG}$
DNA37140	3363	TACCTTTGGAAAAGCTCTCAGGAGACCTCACCTAGATGCCTATTCAAGCTTTGGACAGCC
AX098383	3001	TACCTTTGGAAAAGCTCTCAGGAGACCTCACCTAGATGCCTATTCAAGCTTTGGACAGCC
DNA37140	3423	ATCAGATTGTCAGCCAAGAGCCTTTTATTTGAAAGCTCATTCTTCCCCAGACTTGGACTC
AX098383	3061	ATCAGATTGTCAGCCAAGAGCCTTTTATTTGAAAGCTCATTCTTCCCCAGACTTGGACTC
DNA37140	3483	TGGGTCAGAGGAAGAAGGAAGAAGGACAGATTTTCAGGAAGAAAATCACATTTGTAC
AX098383	3121	TGGGTCAGAGGAAGAAGGAAGAAAGGACAGATTTTCAGGAAGAAAATCACATTTGTAC
DNA37140	3543	CTTTAAACAGACTTTAGAAAACTACAGGACTCCAAATTTTCAGTCTTATGACTTGGACAC
AX098383	3181	CTTTAAACAGACTTTAGAAAACTACAGGACTCCAAATTTTCAGTCTTATGACTTGGACAC
DNA37140	3603	ATAGACTGAATGAGACCAAAGGAAAAGCTTAACATACTACCTCAAGTGAACTTTTATTTA
AX098383	3241	ATAGACTGAATGAGACCAAAGGAAAAGCTTAACATACTACCTCAAGTGAACTTTTATTTA
DNA37140	3663	AAAGAGAGAATCTTATGTTTTTTAAATGGAGTTATGAATTTTAAAAGGATAAAAATGC
AX098383	3301	AAAGAGAGAGAATCTTATGTTTTTTAAATGGAGTTATGAATTTTAAAAGGATAAAAATGC
DNA37140	3723	TTTATTTATACAGATGAACCAAAATTACAAAAAGTTATGAAAATTTTTATACTGGGAATG

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AX098383	3361	TTTATTTATACAGATGAACCAAAATTACAAAAAGTTATGAAAATTTTTATACTGGGAATG
DNA37140	3783	ATGCTCATATAAGAATACCTTTTTAAACTATTTTTTAACTTTGTTTTATGCAAAAAGTA
AX098383	3421	ATGCTCATATAAGAATACCTTTTTAAACTATTTTTTAACTTTGTTTTATGCAAAAAAGTA
DNA37140	3843	TCTTACGTAAATTAATGATATAAATCATGATTATTTTATGTATTTTTATAATGCCAGATT **********************************
AX098383	3481	TCTTACGTAAATTAATGATATAAATCATGATTATTTTATGTATTTTTATAATGCCAGATT
DNA37140	3903	TCTTTTTATGGAAAATGAGTTACTAAAGCATTTTAAATAATACCTGCCTTGTACCATTTT
AX098383	3541	TCTTTTTATGGAAAATGAGTTACTAAAGCATTTTAAATAATACCTGCCTTGTACCATTTT
DNA37140	3963	TTAAATAGAAGTTACTTCATTATATTTTGCACATTATATTTAATAAAATGTGTCAATTTG
AX098383	3601	TTAAATAGAAGTTACTTCATTATATTTTGCACATTATATTTAATAAAATGTGTCAATTTG
DNA37140	4023	AA **
AX098383	3661	AA

```
DNA37140
         3481 TCTGGGTCAGAGGAAGATGGGAAAGAAGGACAGATTTTCAGGAAGAAAATCACATTTGT
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         3481 TCTGGGTCAGAGGAAGATGGGAAAGAAGGACAGATTTTCAGGAAGAAAATCACATTTGT
 AX098387
         3541 ACCTTTAAACAGACTTTAGAAAACTACAGGACTCCAAATTTTCAGTCTTATGACTTGGAC
 DNA37140
             AX098387
         3541 ACCTTTAAACAGACTTTAGAAAACTACAGGACTCCAAATTTTCAGTCTTATGACTTGGAC
 DNA37140
         3601 ACATAGACTGAATGAGACCAAAGGAAAAGCTTAACATACTACCTCAAGTGAACTTTTATT
             ******************
 AX098387
         3601 ACATAGACTGAATGAGACCAAAGGAAAAGCTTAACATACTACCTCAAGTGAACTTTTATT
 DNA37140
         3661 TAAAAGAGAGAGAATCTTATGTTTTTTAAATGGAGTTATGAATTTTAAAAGGATAAAAAT
             *********
 AX098387
         3661 TAAAAGAGAGAGAATCTTATGTTTTTTAAATGGAGTTATGAATTTTAAAAGGATAAAAAT
 DNA37140
         3721 GCTTTATTTATACAGATGAACCAAAATTACAAAAAGTTATGAAAATTTTTATACTGGGAA
             *******************
 AX098387
         3721 GCTTTATTTATACAGATGAACCAAAATTACAAAAAGTTATGAAAATTTTTATACTGGGAA
 DNA37140
         3781 TGATGCTCATATAAGAATACCTTTTTAAACTATTTTTTAACTTTTGTTTTATGCAAAAAG
             ******************
         3781 TGATGCTCATATAAGAATACCTTTTTAAACTATTTTTTAACTTTTTTATGCAAAAAAG
 AX098387
 DNA37140
         3841 TATCTTACGTAAATTAATGATATAAATCATGATTATTTTATGTATTTTTATAATGCCAGA
 AX098387
         3841 TATCTTACGTAAATTAATGATATAAATCATGATTATTTTATGTATTTTATAATGCCAGA
         3901 TTTCTTTTTATGGAAAATGAGTTACTAAAGCATTTTAAATAATACCTGCCTTGTACCATT
 DNA37140
             ******************
 AX098387
         3901 TTTCTTTTTATGGAAAATGAGTTACTAAAGCATTTTAAATAATACCTGCCTTGTACCATT
         3961 TTTTAAATAGAAGTTACTTCATTATATTTTGCACATTATATTTAATAAAATGTGTCAATT
 DNA37140
             AX098387
         3961 TTTTAAATAGAAGTTACTTCATTATATTTTGCACATTATATTTAATAAAATGTGTCAATT
 DNA37140
         4021 TGAAAAAAAAAAAAAAAAAAAAAAAAAAAA
             AX098387
         4021 TGAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAA
>5 AX056647 Sequence 3 from Patent W00075316. (4053 bp) [1 seq]
Score = 4053 (8034 bits), Expect = 0.0
Identities = 4053/4053 (100%), at 1,1-4053,4053, Strand +/+
 DNA37140
           1 AGCCGACGCTGCTCAAGCTGCAACTCTGTTGCAGTTGGCAGTTCTTTTCGGTTTCCCTCC
             AX056647
           1 AGCCGACGCTGCTCAAGCTGCAACTCTGTTGCAGTTGGCAGTTCTTTTCGGTTTCCCTCC
 DNA37140
          61 TGCTGTTTGGGGGCATGAAAGGGCTTCGCCGCCGGGAGTAAAAGAAGGAATTGACCGGGC
            *****************
 AX056647
          61 TGCTGTTTGGGGGCATGAAAGGGCTTCGCCGCCGGGAGTAAAAGAAGGAATTGACCGGGC
 DNA37140
         ***********
 AX056647
```

181 TTGTGCCGGGCCCCGAGCGCGCCGGCTGGGAGCTTCGGGTAGAGACCTAGGCCGCTGG

DNA37140

		*****************
AX056647	181	TTGTGCCGGGCCCGAGCGCGCGCGGCTGGGAGCTTCGGGTAGAGACCTAGGCCGCTGG
DNA3714G	241	ACCGCGATGAGCGCGCGAGCCTCCGTGCGCGCGCGCGGGGTTGGGGCTGCTGTGC **********
AX056647	241	ACCGCGATGAGCGCGCGAGCCTCCGTGCGCGCGCGCGGGGTTGGGGGCTGCTGTGC
DNA37140	301	GCGGTGCTGGGGCGCTGGCCGGTCCGACAGCGGCGGTCGCGGGGAACTCGGGCAGCCC
AX056647	301	GCGGTGCTGGGCGCCGCTCCGACAGCGGCGGTCGCGGGGAACTCGGGCAGCCC
DNA37140	361	TCTGGGGTAGCCGCCGAGCGCCCATGCCCCACTACCTGCCGCTGCCTCGGGGACCTGCTG ********************************
AX056647	361	TCTGGGGTAGCCGCCGAGCGCCCATGCCCCACTACCTGCCGCTGCCTCGGGGACCTGCTG
DNA37140	421	GACTGCAGTCGTAAGCGGCTAGCGCGTCTTCCCGAGCCACTCCCGTCCTGGGTCGCTCGG
AX056647	421	GACTGCAGTCGTAAGCGGCTAGCGCGTCTTCCCGAGCCACTCCCGTCCTGGGTCGCTCGG
DNA37140	481	CTGGACTTAAGTCACAACAGATTATCTTTCATCAAGGCAAGTTCCATGAGCCACCTTCAA
AX056647	481	CTGGACTTAAGTCACAACAGATTATCTTTCATCAAGGCAAGTTCCATGAGCCACCTTCAA
DNA37140	541	AGCCTTCGAGAAGTGAAACTGAACAATGAATTGGAGACCATTCCAAATCTGGGACCA ********************************
AX056647	541	AGCCTTCGAGAAGTGAAACTGAACAATGAATTGGAGACCATTCCAAATCTGGGACCA
DNA37140	601	GTCTCGGCAAATATTACACTTCTCCTTGGCTGGAAACAGGATTGTTGAAATACTCCCT
AX056647	601	GTCTCGGCAAATATTACACTTCTCCCTTGGCTGGAAACAGGATTGTTGAAATACTCCCT
DNA37140	661	GAACATCTGAAAGAGTTTCAG <sup>†</sup> CCCTTGAAACTTTGGACCTTAGCAGCAACAATATTTCA **************************
AX056647	661	GAACATCTGAAAGAGTTTCAGTCCCTTGAAACTTTGGACCTTAGCAGCAACAATATTTCA
DNA37140	721	GAGCTCCAAACTGCATTTCCAGCCCTACAGCTCAAATATCTGTATCTCAACAGCAACCGA **************************
AX056647	721	GAGCTCCAAACTGCATTTCCAGCCCTACAGCTCAAATATCTGTATCTCAACAGCAACCGA
DNA37140	781	GTCACATCAATGGAACCTGGGTATTTTGACAATTTGGCCAACACACTCCTTGTGTTAAAG ****************************
AX056647	781	GTCACATCAATGGAACCTGGGTATTTTGACAATTTGGCCAACACACTCCTTGTGTTAAAG
DNA37140	841	CTGAACAGGAACCGAATCTCAGCTATCCCACCCAAGATGTTTAAACTGCCCCAACTGCAA ***********************************
AX056647	841	CTGAACAGGAACCGAATCTCAGCTATCCCACCCAAGATGTTTAAACTGCCCCAACTGCAA
DNA37140	901	CATCTCGAATTGAACCGAAACAAGATTAAAAATGTAGATGGACTGACATTCCAAGGCCTT *********************************
AX056647	901	CATCTCGAATTGAACCGAAACAAGATTAAAAATGTAGATGGACTGACATTCCAAGGCCTT
DNA37140		GGTGCTCTGAAGTCTCTGAAAATGCAAAGAAATGGAGTAACGAAACTTATGGATGG
AX056647	961	GGTGCTCTGAAGTCTCTGAAAATGCAAAGAAATGGAGTAACGAAACTTATGGATGG
DNA37140		TTTTGGGGGCTGAGCAACATGGAAATTTTGCAGCTGGACCATAACAACCTAACAGAGATT ********************************

AX056647	1021	TTTTGGGGGCTGAGCAACATGGAAATTTTGCAGCTGGACCATAACAACCTAACAGAGATT
DNA37140	1081	ACCAAAGGCTGGCTTTACGGCTTGCTGATGCTGCAGGAACTTCATCTCAGCCAAAATGCC
AX056647	1081	ACCAAAGGCTGGCTTTACGGCTTGCTGATGCTGCAGGAACTTCATCTCAGCCAAAATGCC
DNA37140	1141	ATCAACAGGATCAGCCCTGATGCCTGGGAGTTCTGCCAGAAGCTCAGTGAGCTGGACCTA
AX056647	1141	ATCAACAGGATCAGCCCTGATGCCTGGGAGTTCTGCCAGAAGCTCAGTGAGCTGGACCTA
DNA37140	1201	ACTTTCAATCACTTATCAAGGTTAGATGATTCAAGCTTCCTTGGCCTAAGCTTACTAAAT
AX056647	1201	ACTTTCAATCACTTATCAAGGTTAGATGATTCAAGCTTCCTTGGCCTAAGCTTACTAAAT
DNA37140	1261	ACACTGCACATTGGGAACAACAGAGTCAGCTACATTGCTGATTGTGCCTTCCGGGGGCTT
AX056647	1261	ACACTGCACATTGGGAACAACAGAGTCAGCTACATTGCTGATTGTGCCTTCCGGGGGCTT
DNA37140	1321	TCCAGTTTAAAGACTTTGGATCTGAAGAACAATGAAATTTCCTGGACTATTGAAGACATG
AX056647	1321	TCCAGTTTAAAGACTTTGGATCTGAAGAACAATGAAATTTCCTGGACTATTGAAGACATG
DNA37140	1381	AATGGTGCTTTCTCTGGGCTTGACAAACTGAGGCGACTGATACTCCAAGGAAATCGGATC
AX056647	1381	AATGGTGCTTTCTCTGGGCTTGACAAACTGAGGCGACTGATACTCCAAGGAAATCGGATC
DNA37140	1441	CGTTCTATTACTAAAAAAGCCTTCACTGGTTTGGATGCATTGGAGCATCTAGACCTGAGT
AX056647	1441	CGTTCTATTACTAAAAAAGCCTTCACTGGTTTGGATGCATTGGAGCATCTAGACCTGAGT
DNA37140	1501	GACAACGCAATCATGTCTTTAÇAAGGCAATGCATTTTCACAAATGAAGAAACTGCAACAA
AX056647	1501	GACAACGCAATCATGTCTTTACAAGGCAATGCATTTTCACAAATGAAGAAACTGCAACAA
DNA37140	1561	TTGCATTTAAATACATCAAGCCTTTTGTGCGATTGCCAGCTAAAATGGCTCCCACAGTGG
AX056647	1561	TTGCATTTAAATACATCAAGCCTTTTGTGCGATTGCCAGCTAAAATGGCTCCCACAGTGG
DNA37140	1621	GTGGCGGAAAACAACTTTCAGAGCTTTGTAAATGCCAGTTGTGCCCATCCTCAGCTGCTA ************************************
AX056647	1621	GTGGCGGAAAACAACTTTCAGAGCTTTGTAAATGCCAGTTGTGCCCATCCTCAGCTGCTA
DNA37140	1681	AAAGGAAGAAGCATTTTTGCTGTTAGCCCAGATGGCTTTGTGTGTG
AX056647	1681	AAAGGAAGAAGCATTTTTGCTGTTAGCCCAGATGGCTTTGTGTGTG
DNA37140	1741	CCCCAGATCACGGTTCAGCCAGAAACACAGTCGGCAATAAAAGGTTCCAATTTGAGTTTC
AX056647	1741	CCCCAGATCACGGTTCAGCCAGAAACACAGTCGGCAATAAAAGGTTCCAATTTGAGTTTC
DNA37140	1801	ATCTGCTCAGCTGCCAGCAGCAGTGATTCCCCAATGACTTTTGCTTGGAAAAAAAGACAAT
AX056647	1801	ATCTGCTCAGCTGCCAGCAGCAGTGATTCCCCCAATGACTTTTGCTTGGAAAAAAAGACAAT
DNA37140	1861	GAACTACTGCATGATGCTGAAATGGAAAATTATGCACACCTCCGGGCCCAAGGTGGCGAG
AX056647	1861	GAACTACTGCATGATGCTGAAATGGAAAATTATGCACACCTCCGGGCCCAAGGTGGCGAG

DNA37140	1921	GTGATGGAGTATACCACCATCCTTCGGCTGCGCGAGGTGGAATTTGCCAGTGAGGGGAAA ***************************
AX056647	1921	GTGATGGAGTATACCACCATCCTTCGGCTGCGCGAGGTGGAATTTGCCAGTGAGGGGAAA
DNA37140	1981	TATCAGTGTGTCATCTCCAATCACTTTGGTTCATCCTACTCTGTCAAAGCCAAGCTTACA *********************************
AX056647	1981	TATCAGTGTCTCCAATCACTTTGGTTCATCCTACTCTGTCAAAGCCAAGCTTACA
DNA37140	2041	GTAAATATGCTTCCCTCATTCACCAAGACCCCCATGGATCTCACCATCCGAGCTGGGGCC ******************************
AX056647	2041	GTAAATATGCTTCCCTCATTCACCAAGACCCCCATGGATCTCACCATCCGAGCTGGGGCC
DNA37140	2101	ATGGCACGCTTGGAGTGTGCTGCTGTGGGGCACCCAGCCCCCAGATAGCCTGGCAGAAG *******************************
AX056647	2101	ATGGCACGCTTGGAGTGTGCTGCTGTGGGGCACCCAGCCCCCAGATAGCCTGGCAGAAG
DNA37140	2161	GATGGGGGCACAGACTTCCCAGCTGCACGGGAGAGACGCATGCAT
AX056647	2161	GATGGGGGCACAGACTTCCCAGCTGCACGGGAGAGACGCATGCAT
DNA37140	2221	GACGTGTTCTTTATCGTGGATGTGAAGATAGAGGACATTGGGGTATACAGCTGCACAGCT
AX056647	2221	GACGTGTTCTTTATCGTGGATGTGAAGATAGAGGACATTGGGGTATACAGCTGCACAGCT
DNA37140	2281	CAGAACAGTGCAGGAAGTATTTCAGCAAATGCAACTCTGACTGTCCTAGAAACACCATCA
AX056647	2281	CAGAACAGTGCAGGAAGTATTTCAGCAAATGCAACTCTGACTGTCCTAGAAACACCATCA
DNA37140	2341	TTTTTGCGGCCACTGTTGGACCGAACTGTAACCAAGGGAGAAACAGCCGTCCTACAGTGC ***********************************
AX056647	2341	TTTTTGCGGCCACTGTTGGACCGAACTGTAACCAAGGGAGAAACAGCCGTCCTACAGTGC
DNA37140	2401	ATTGCTGGAGGAAGCCCTCCCCCTAAACTGAACTGGACCAAAGATGATAGCCCATTGGTG ********************************
AX056647	2401	ATTGCTGGAGGAAGCCCTCCCCCTAAACTGAACTGGACCAAAGATGATAGCCCATTGGTG
DNA37140	2461	GTAACCGAGAGGCACTTTTTTGCAGCAGGCAATCAGCTTCTGATTATTGTGGACTCAGAT ***********************************
AX056647	2461	GTAACCGAGAGGCACTTTTTTGCAGCAGGCAATCAGCTTCTGATTATTGTGGACTCAGAT
DNA37140	2521	GTCAGTGATGCTGGGAAATACACATGTGAGATGTCTAACACCCTTGGCACTGAGAGAGA
AX056647	2521	GTCAGTGATGCTGGGAAATACACATGTGAGATGTCTAACACCCTTGGCACTGAGAGAGGA
DNA37140	2581	AACGTGCGCCTCAGTGTGATCCCCACTCCAACCTGCGACTCCCCTCAGATGACAGCCCCA *****************************
AX056647	2581	AACGTGCGCCTCAGTGTGATCCCCACTCCAACCTGCGACTCCCCTCAGATGACAGCCCCA
DNA37140	2641	TCGTTAGACGATGACGGATGGGCCACTGTGGGTGTCGTGATCATAGCCGTGGTTTGCTGT ************************
AX056647	2641	TCGTTAGACGATGACGGATGGGCCACTGTGGGTGTCGTGATCATAGCCGTGGTTTGCTGT
DNA37140	2701	GTGGTGGCACGTCACTCGTGTGGGTGGTCATCATATACCACACAAGGCGGAGGAATGAA
AX056647	2701	GTGGTGGGCACGTCACTCGTGTGGGTGGTCATCATATACCACACAAGGCGGAGGAATGAA

DNA37140	2761	GATTGCAGCATTACCAACACAGATGAGACCAACTTGCCAGCAGATATTCCTAGTTATTTG
AX056647	2761	GATTGCAGCATTACCAACACAGATGAGACCAACTTGCCAGCAGATATTCCTAGTTATTTG
DNA37140	2821	TCATCTCAGGGAACGTTAGCTGACAGGCAGGATGGGTACGTGTCTTCAGAAAGTGGAAGC
AX056647	2821	TCATCTCAGGGAACGTTAGCTGACAGGCAGGATGGGTACGTGTCTTCAGAAAGTGGAAGC
DNA37140	2881	CACCACCAGTTTGTCACATCTTCAGGTGCTGGATTTTTCTTACCACAACATGACAGTAGT
AX056647	2881	CACCACCAGTTTGTCACATCTTCAGGTGCTGGATTTTTCTTACCACAACATGACAGTAGT
DNA37140	2941	GGGACCTGCCATATTGACAATAGCAGTGAAGCTGATGTGGAAGCTGCCACAGATCTGTTC ********************************
AX056647	2941	GGGACCTGCCATATTGACAATAGCAGTGAAGCTGATGTGGAAGCTGCCACAGATCTGTTC
DNA37140	3001	CTTTGTCCGTTTTTGGGATCCACAGGCCCTATGTATTTGAAGGGAAATGTGTATGGCTCA
AX056647	3001	CTTTGTCCGTTTTTGGGATCCACAGGCCCTATGTATTTGAAGGGAAATGTGTATGGCTCA
DNA37140	3061	GATCCTTTTGAAACATATCATACAGGTTGCAGTCCTGACCCAAGAACAGTTTTAATGGAC
AX056647	3061	GATCCTTTTGAAACATATCATACAGGTTGCAGTCCTGACCCAAGAACAGTTTTAATGGAC
DNA37140	3121	CACTATGAGCCCAGTTACATAAAGAAAAAGGAGTGCTACCCATGTTCTCATCCTTCAGAA
AX056647	3121	CACTATGAGCCCAGTTACATAAAGAAAAAGGAGTGCTACCCATGTTCTCATCCTTCAGAA
DNA37140	3181	GAATCCTGCGAACGGAGCTTCAGTAATATATCGTGGCCTTCACATGTGAGGAAGCTACTT
AX056647	3181	${\tt GAATCCTGCGAACGGAGCTTCAGTAATATCGTGGCCTTCACATGTGAGGAAGCTACTT}.$
DNA37140	3241	AACACTAGTTACTCTCACAATGAAGGACCTGGAATGAAAAATCTGTGTCTAAACAAGTCC ***********************************
AX056647	3241	AACACTAGTTACTCTCACAATGAAGGACCTGGAATGAAAAATCTGTGTCTAAACAAGTCC
DNA37140	3301	TCTTTAGATTTTAGTGCAAATCCAGAGCCAGCGTCGGTTGCCTCGAGTAATTCTTTCATG ************************************
AX056647	3301	TCTTTAGATTTTAGTGCAAATCCAGAGCCAGCGTCGGTTGCCTCGAGTAATTCTTTCATG
DNA37140	3361	GGTACCTTTGGAAAAGCTCTCAGGAGACCTCACCTAGATGCCTATTCAAGCTTTGGACAG *********************************
AX056647	3361	GGTACCTTTGGAAAAGCTCTCAGGAGACCTCACCTAGATGCCTATTCAAGCTTTGGACAG
DNA37140	3421	CCATCAGATTGTCAGCCAAGAGCCTTTTATTTGAAAGCTCATTCTTCCCCAGACTTGGAC
AX056647	3421	CCATCAGATTGTCAGCCAAGAGCCTTTTATTTGAAAGCTCATTCTTCCCCAGACTTGGAC
DNA37140	3481	TCTGGGTCAGAGGAAGATGGGAAAGAAAGGACAGATTTTCAGGAAGAAAATCACATTTGT *********************************
AX056647	3481	TCTGGGTCAGAGGAAGATGGGAAAGAAAGGACAGATTTTCAGGAAGAAAATCACATTTGT
DNA37140	3541	ACCTTTAAACAGACTTTAGAAAACTACAGGACTCCAAATTTTCAGTCTTATGACTTGGAC
AX056647	3541	ACCTTTAAACAGACTTTAGAAAACTACAGGACTCCAAATTTTCAGTCTTATGACTTGGAC
DNA37140	3601	ACATAGACTGAATGAGACCAAAGGAAAAGCTTAACATACTACCTCAAGTGAACTTTTATT

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	AX056647	3601	ACATAGACTGAATGAGACCAAAGGAAAAGCTTAACATACTACCTCAAGTGAACTTTTATT
	DNA37140	3661	TAAAAGAGAGAGAATCTTATGTTTTTTAAATGGAGTTATGAATTTTAAAAGGATAAAAAT
	AX056647	3661	TAAAAGAGAGAGAATCTTATGTTTTTTAAATGGAGTTATGAATTTTAAAAGGATAAAAAT
	DNA37140	3721	GCTTTATTTATACAGATGAACCAAAATTACAAAAAGTTATGAAAATTTTTATACTGGGAA
	AX056647	3721	GCTTTATTTATACAGATGAACCAAAATTACAAAAAGTTATGAAAATTTTTATACTGGGAA
	DNA37140	3781	TGATGCTCATATAAGAATACCTTTTTAAACTATTTTTTAACTTTTGTTTTATGCAAAAAAG
	AX056647	3781	TGATGCTCATATAAGAATACCTTTTTAAACTATTTTTTAACTTTGTTTTATGCAAAAAAG
	DNA37140	3841	TATCTTACGTAAATTAATGATATAAATCATGATTATTTTATGTATTTTTATAATGCCAGA
	AX056647	3841	TATCTTACGTAAATTAATGATATAAATCATGATTATTTTATGTATTTTTATAATGCCAGA
	DNA37140	3901	TTTCTTTTTATGGAAAATGAGTTACTAAAGCATTTTAAATAATACCTGCCTTGTACCATT
	AX056647	3901	TTTCTTTTTATGGAAAATGAGTTACTAAAGCATTTTAAATAATACCTGCCTTGTACCATT
	DNA37140	3961	TTTTAAATAGAAGTTACTTCATTATATTTTGCACATTATATTTAATAAAATGTGTCAATT *********************************
	AX056647	3961	TTTTAAATAGAAGTTACTTCATTATATTTTGCACATTATATTTAATAAAATGTGTCAATT
	DNA37140	4021	TGAAAAAAAAAAAAAAAAAAAAAAAAA
			************
	AX056647	4021	**************************************
>6	P_AAC8442	22 Hun	TGAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAA
>6 (4	P_AAC8442 053 bp) [1	22 Hun seg]	TGAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAA
>6 (4 S	P_AAC8442 053 bp) [1 core = 402	22 Hum seg]	TGAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAA
>6 (4 S I	P_AAC8442 053 bp) [1 core = 402	22 Hum seg] 21 (79 = 404	TGAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAA
>6 (4 S I	P_AAC8442 053 bp) [1 core = 402 dentities	22 Hum seg] 21 (79 = 404	TGAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAA
>6 (4 S I	P_AAC8442 053 bp) [1 core = 402 dentities DNA37140	22 Hum seg] 21 (79 = 404	TGAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAA
>6 (4 S I	P_AAC8442 053 bp) [1 core = 402 dentities DNA37140 AAC84422	22 Hum seg] 21 (79 = 404 1	TGAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAA
>6 (4 S I	P_AAC8442 053 bp) [1 core = 402 dentities DNA37140 AAC84422 DNA37140	22 Hum seg] 21 (79 = 404 1 1 61	TGAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAA
>6 (4 S I	P_AAC84422 053 bp) [1 core = 402 dentities DNA37140 AAC84422 DNA37140 AAC84422	22 Hum seg] 21 (79 = 404 1 1 61 61	TGAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAA
>6 (4 S I P	P_AAC84422 053 bp) [1 core = 402 dentities  DNA37140  AAC84422  DNA37140  AAC84422  DNA37140	22 Hum . seg] 21 (79 = 404  1  61  121	TGAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAA
>6 (4 S I P	P_AAC84422 053 bp) [1 core = 402 dentities  DNA37140  AAC84422  DNA37140  AAC84422  DNA37140  AAC84422	22 Hum. seg] 21 (79 = 404  1  61  121  121  181	TGAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAA
>6 (4 S I P	P_AAC8442 053 bp) [1 core = 402 dentities DNA37140 AAC84422 DNA37140 AAC84422 DNA37140 AAC84422 DNA37140	22 Hum seg] 21 (79 = 404  1 61 121 181	TGAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAA
>6 (4 S I P	P_AAC84422 053 bp) [1 core = 402 dentities  DNA37140  AAC84422  DNA37140  AAC84422  DNA37140  AAC84422  DNA37140  AAC84422  AAC84422	22 Hum seg] 21 (79 = 404  1  61  121  181  181  241	TGAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAA

P_AAC84422	301	GCGGTGCTGGGCGCCGCTGGCCGGTCCGACAGCGGCGGTCGCGGGGAACTCGGGCAGCCC
DNA37140	361	TCTGGGGTAGCCGCCGAGCGCCCATGCCCACTACCTGCCGCTGCCTCGGGGACCTGCTG ********************************
P_AAC84422	361	TCTGGGGTAGCCGCCGAGCGCCCATGCCCCACTACCTGCCGCTGCCTCGGGGACCTGCTG
DNA37140	421	GACTGCAGTCGTAAGCGGCTAGCGCGTCTTCCCGAGCCACTCCCGTCCTGGGTCGCTCGG
P_AAC84422	421	${\tt GACTGCAGTCGTAAGCGGCTAGCGCGTCTTCCCGAGCCACTCCCGTCCTGGGTCGCTCGGGTCGGGGTCGGGTCGGGTCGGGGTCGGGGTCGGGGGTCGGGGGG$
DNA37140	481	CTGGACTTAAGTCACAACAGATTATCTTTCATCAAGGCAAGTTCCATGAGCCACCTTCAA
P_AAC84422	481	$\tt CTGGACTTAAGTCACAACAGATTATCTTTCATCAAGGCAAGTTCCATGAGCCACCTTCAAGGCAAGTTCCATGAGCCACCTTCAAGGCAAGTTCCATGAGCCACCTTCAAGGCAAGTTCCATGAGCCACCTTCAAGGCAAGTTCCATGAGCCACCTTCAAGGCAAGTTCCATGAGCCACCTTCAAGGCAAGTTCCATGAGCCACCTTCAAGGCAAGTTCCATGAGCCACCTTCAAGGCAAGTTCCATGAGCCACCTTCAAGGCAAGTTCCATGAGCCACCTTCAAGGCAAGTTCCATGAGCCACCTTCAAGGCAAGTTCCATGAGCCACCTTCAAGGCAAGTTCCATGAGCCACCTTCAAGGCAAGTTCCATGAGCCACCTTCAAGGCCACCTTCAAGGCAAGTTCCATGAGCCACCTTCAAGGCAAGTTCCATGAGCCACCTTCAAGGCCACCTTCAAGGCCACCTTCAAGGCCACCTTCAAGGCCACCTTCAAGGCCACCTTCAAGGCCACCTTCAAGGCCACGTTCAAGGCCACCTTCAAGGCCAAGTTCCATGAGGCAAGTTCCATGAGGCAAGTTCCATGAGGCAAGTTCCATGAGGCAAGTTCCATGAGGCAAGTTCCATGAGGCAAGGTTCCATGAGGCAAGTTCCATGAGGCAAGTTCCATGAGGCAAGTTCCATGAGGCAAGTTCCATGAGGCAAGTTCCATGAGAGGCAAGTTCCATGAGAGGCAAGTTCCATGAGGCAAGTTCCATGAGGCAAGTTCCATGAGGCAAGTTCCATGAGGCAAGTTCCATGAGAGGCAAGTTCCATGAGGCAAGTTCCATGAGGCAAGTTCCATGAGGCAAGTTCCATGAGGCAAGTTCCATGAGGCAAGTTCCATGAGGCAAGTTCCATGAGGCAAGTTCCATGAGGCAAGTTCCATGAGGCAAGTTCCATGAGGCAAGTTCCATGAGGCAAGTTCCATGAGGCAAGTTCCATGAGAGGCAAGTTCCATGAGAGGCAAGTTCAAGAGGCAAGTTCAAGAGGCAAGTTCAAGAGGCAAGTTCAAGAGAGAG$
DNA37140	541	AGCCTTCGAGAAGTGAAACTGAACAACAATGAATTGGAGACCATTCCAAATCTGGGACCA ********************************
P_AAC84422	541	${\tt AGCCTTCGAGAAGTGAAACTGAACAACAATGAATTGGAGACCATTCCAAATCTGGGACCAACCA$
DNA37140	601	GTCTCGGCAAATATTACACTTCTCTCTTGGCTGGAAACAGGATTGTTGAAATACTCCCT
P_AAC84422	601	$\tt GTCTCGGCAAATATTACACTTCTCTCTTGGCTGGAAACAGGATTGTTGAAATACTCCCTTGGCTGGAAACAGGATTGTTGAAATACTCCCTTGGCTGGAAACAGGATTGTTGAAATACTCCCTTGGCTGGAAACAGGATTGTTGAAATACTCCCTTGGCTGGAAACAGGATTGTTGAAATACTCCCTTGGCTGGAAACAGGATTGTTGAAATACTCCCTTGGCTGGAAACAGGATTGTTGAAATACTCCCTTGGCTGGAAACAGGATTGTTGAAATACTCCCTTGGCTGGAAACAGGATTGTTGAAAATACTCCCTTGGCTGGAAACAGGATTGTTGAAAATACTCCCTTGGCTGGAAACAGGATTGTTGAAAATACTCCCTTGGCTGGAAACAGGATTGTTGAAAATACTCCCTTGGCTGGAAACAGGAATTGTTGAAAATACTCCCTTGGCTGGAAACAGGAATTGTTGAAAATACTCCCTTGGCTGGAAACAGGAATTGTTGAAAATACTCCCTTGGCTGGAAACAGGAATTGTTGAAAATACTCCCTTGGCTGGAAACAGGAATTGTTGAAAATACTCCCTTGGCTGAAACAGGAATTGTTGAAAATACTCCCTTGGCTGAAAACAGGAATTGTTGAAAATACTCCCTTGGCTGAAAATACTCCCTTGGCTGAAAATACTTCCCTTGGCTGAAAATACTTCCCTTGGCTGAAAATACTTCCCTTGGCTGAAAATACTTCCCTTGGCTGAAAATACTTCCCTTGGCTGAAAATACTTCCCTTGGCTGAAAAAATACTTCCCTTGGCTGAAAAAAAA$
DNA37140	661	GAACATCTGAAAGAGTTTCAGTCCCTTGAAACTTTGGACCTTAGCAGCAACAATATTTCA
P_AAC84422	661	GAACATCTGAAAGAGTTTCAGTCCCTTGAAACTTTGGACCTTAGCAGCAACAATATTTCA
DNA37140	721	GAGCTCCAAACTGCATTTCCAGCCCTACAGCTCAAATATCTGTATCTCAACAGCAACCGA
P_AAC84422	721	GAGCTCCAAACTGCATTTCCAGCCCTACAGCTCAAATATCTGTATCTCAACAGCAACCGA
DNA37140	781	GTCACATCAATGGAACCTGGGTATTTTGACAATTTGGCCAACACACTCCTTGTGTTAAAG
P_AAC84422	781	$\tt GTCACATCAATGGAACCTGGGTATTTTGACAATTTGGCCAACACACTCCTTGTGTTAAAGGACACACAC$
DNA37140	841	CTGAACAGGAACCGAATCTCAGCTATCCCACCCAAGATGTTTAAACTGCCCCAACTGCAA
P_AAC84422	841	$\tt CTGAACAGGAACCGAATCTCAGCTATCCCACCCAAGATGTTTAAACTGCCCCAACTGCAACTAACAACTGCAACTGCAACTGCAACTGCAACTGCAACTAACAACTAACAACAACAACAACAACAACAACAACA$
DNA37140	901	CATCTCGAATTGAACCGAAACAAGATTAAAAATGTAGATGGACTGACATTCCAAGGCCTT
P_AAC84422	901	CATCTCGAATTGAACCGAAACAAGATTAAAAATGTAGATGGACTGACATTCCAAGGCCTT
DNA37140	961	GGTGCTCTGAAGTCTCTGAAAATGCAAAGAAATGGAGTAACGAAACTTATGGATGG
P_AAC84422	961	GGTGCTCTGAAGATGCAAAGAAATGGAGTAACGAAACTTATGGATGG
DNA37140	1021	TTTTGGGGGCTGAGCAACATGGAAATTTTGCAGCTGGACCATAACAACCTAACAGAGATT
P_AAC84422	1021	TTTTGGGGGCTGAGCAACATGGAAATTTTGCAGCTGGACCATAACAACCTAACAGAGATT
DNA37140	1081	ACCAAAGGCTGGCTTTACGGCTTGCTGATGCTGCAGGAACTTCATCTCAGCCAAAATGCC
P_AAC84422	1081	ACCAAAGGCTGGCTTTACGGCTTGCTGATGCTGCAGGAACTTCATCTCAGCCAAAATGCC
DNA37140	1141	ATCAACAGGATCAGCCCTGATGCCTGGGAGTTCTGCCAGAAGCTCAGTGAGCTGGACCTA

P_AAC84422	1141	ATCAACAGGATCAGCCCTGATGCCTGGGAGTTCTGCCAGAAGCTCAGTGAGCTGGACCTA
DNA37140		ACTTTCAATCACTTATCAAGGTTAGATGATTCAAGCTTCCTTGGCCTAAGCTTACTAAAT *****************************
P_AAC84422	1201	ACTTTCAATCACTTATCAAGGTTAGATGATTCAAGCTTCCTTGGCCTAAGCTTACTAAAT
DNA37140	1261	ACACTGCACATTGGGAACAACAGAGTCAGCTACATTGCTGATTGTGCCTTCCGGGGGCTT ********************
P_AAC84422	1261	ACACTGCACATTGGGAACAACAGAGTCAGCTACATTGCTGATTGTGCCTTCCGGGGGCTT
DNA37140	1321	TCCAGTTTAAAGACTTTGGATCTGAAGAACAATGAAATTTCCTGGACTATTGAAGACATG
P_AAC84422	1321	TCCAGTTTAAAGACTTTGGATCTGAAGAACAATGAAATTTCCTGGACTATTGAAGACATG
DNA37140	1381	AATGGTGCTTTCTCTGGGCTTGACAAACTGAGGCGACTGATACTCCAAGGAAATCGGATC ************************************
P_AAC84422	1381	AATGGTGCTTTCTCTGGGCTTGACAAACTGAGGCGACTGATACTCCAAGGAAATCGGATC
DNA37140	1441	CGTTCTATTACTAAAAAAGCCTTCACTGGTTTGGATGCATTGGAGCATCTAGACCTGAGT
P_AAC84422	1441	CGTTCTATTACTAAAAAAGCCTTCACTGGTTTGGATGCATTGGAGCATCTAGACCTGAGT
DNA37140	1501	GACAACGCAATCATGTCTTTACAAGGCAATGCATTTTCACAAATGAAGAAACTGCAACAA
P_AAC84422	1501	GACAACGCAATCATGTCTTTACAAGGCAATGCATTTTCACAAATGAAGAAACTGCAACAA
DNA37140	1561	TTGCATTTAAATACATCAAGCCTTTTGTGCGATTGCCAGCTAAAATGGCTCCCACAGTGG
P_AAC84422	1561	TTGCATTTAAATACATCAAGCCTTTTGTGCGATTGCCAGCTAAAATGGCTCCCACAGTGG
DNA37140	1621	GTGGCGGAAAACAACTTTCAGAGCTTTGTAAATGCCAGTTGTGCCCATCCTCAGCTGCTA
P_AAC84422	1621	GTGGCGGAAAACAACTTTCAGAGCTTTGTAAATGCCAGTTGTGCCCATCCTCAGCTGCTA
DNA37140	1681	AAAGGAAGAAGCATTTTTGCTGTTAGCCCAGATGGCTTTGTGTGTG
P_AAC84422	1681	AAAGGAAGAAGCATTTTTGCTGTTAGCCCAGATGGCTTTGTGTGTG
DNA37140	1741	CCCCAGATCACGGTTCAGCCAGAAACACAGTCGGCAATAAAAGGTTCCAATTTGAGTTTC
P_AAC84422	1741	$\tt CCCCAGATCACGGTTCAGCCAGAAACACAGTCGGCAATAAAAGGTTCCAATTTGAGTTTC$
DNA37140	1801	ATCTGCTCAGCTGCCAGCAGCAGTGATTCCCCAATGACTTTTGCTTGGAAAAAAAGACAAT
P_AAC84422	1801	ATCTGCTCAGCTGCCAGCAGCAGTGATTCCCCAATGACTTTTGCTTGGAAAAAAAGACAAT
DNA37140	1861	GAACTACTGCATGATGCTGAAATGGAAAATTATGCACACCTCCGGGCCCAAGGTGGCGAG
P_AAC84422	1861	GAACTACTGCATGATGCTGAAATGGAAAATTATGCACACCTCCGGGCCCAAGGTGGCGAG
DNA37140	1921	GTGATGGAGTATACCACCATCCTTCGGCTGCGCGAGGTGGAATTTGCCAGTGAGGGGAAA
P_AAC84422	1921	GTGATGGAGTATACCACCATCCTTCGGCTGCGCGAGGTGGAATTTGCCAGTGAGGGGAAA
DNA37140	1981	TATCAGTGTGTCATCTCCAATCACTTTGGTTCATCCTACTCTGTCAAAGCCAAGCTTACA
P_AAC84422	1981	TATCAGTGTCATCTCCAATCACTTTGGTTCATCCTACTCTGTCAAAGCCAAGCTTACA

DNA37140	2041	GTAAATATGCTTCCCTCATTCACCAAGACCCCCATGGATCTCACCATCCGAGCTGGGGCC
P_AAC84422	2041	GTAAATATGCTTCCCTCATTCACCAAGACCCCCATGGATCTCACCATCCGAGCTGGGGCC
DNA37140	2101	ATGGCACGCTTGGAGTGTGCTGCTGTGGGGCACCCAGCCCCCCAGATAGCCTGGCAGAAG
P_AAC84422	2101	ATGGCACGCTTGGAGTGTGCTGCTGTGGGGCACCCAGCCCCCAGATAGCCTGGCAGAAG
DNA37140	2161	GATGGGGGCACAGACTTCCCAGCTGCACGGGAGAGACGCATGCAT
P_AAC84422	2161	GATGGGGGCACAGACTTCCCAGCTGCACGGAGAGACGCATGCAT
DNA37140	2221	GACGTGTTCTTTATCGTGGATGTGAAGATAGAGGACATTGGGGTATACAGCTGCACAGCT
P_AAC84422	2221	GACGTGTTCTTTATCGTGGATGTGAAGATAGAGGACATTGGGGTATACAGCTGCACAGCT
DNA37140	2281	CAGAACAGTGCAGGAAGTATTTCAGCAAATGCAACTCTGACTGTCCTAGAAACACCATCA
P_AAC84422	2281	CAGAACAGTGCAGGAAGTATTTCAGCAAATGCAACTCTGACTGTCCTAGAAACACCATCA
DNA37140	2341	TTTTTGCGGCCACTGTTGGACCGAACTGTAACCAAGGGAGAAACAGCCGTCCTACAGTGC
P_AAC84422	2341	TTTTTGCGGCCACTGTTGGACCGAACTGTAACCAAGGGAGAAACAGCCGTCCTACAGTGC
DNA37140	2401	ATTGCTGGAGGAAGCCCTCCCCCTAAACTGAACTGGACCAAAGATGATAGCCCATTGGTG
P_AAC84422	2401	ATTGCTGGAGGAAGCCCTCCCCCTAAACTGAACTGGACCAAAGATGATAGCCCATTGGTG
DNA37140	2461	GTAACCGAGAGGCACTTTTTTGCAGCAGGCAATCAGCTTCTGATTATTGTGGACTCAGAT ***********************************
P_AAC84422	2461	GTAACCGAGAGGCACTTTTTTGCAGCAGGCAATCAGCTTCTGATTATTGTGGACTCAGAT
DNA37140	2521	GTCAGTGATGCTGGGAAATACACATGTGAGATGTCTAACACCCTTGGCACTGAGAGAGGA
P_AAC84422	2521	GTCAGTGATGCTGGGAAATACACATGTGAGATGTCTAACACCCTTGGCACTGAGAGAGGA
DNA37140	2581	AACGTGCGCCTCAGTGTGATCCCCACTCCAACCTGCGACTCCCCTCAGATGACAGCCCCA
P_AAC84422	2581	AACGTGCGCCTCAGTGTGATCCCCACTCCAACCTGCGACTCCCCTCAGATGACAGCCCCA
DNA37140	2641	TCGTTAGACGATGACGGATGGGCCACTGTGGGTGTCGTGATCATAGCCGTGGTTTGCTGT
P_AAC84422	2641	**************************************
DNA37140	2701	GTGGTGGCACGTCACTCGTGTGGGTGGTCATCATATACCACACAAGGCGGAGGAATGAA
P_AAC84422	2701	**************************************
DNA37140	2761	GATTGCAGCATTACCAACACAGATGAGACCAACTTGCCAGCAGATATTCCTAGTTATTTG
P_AAC84422	2761	**************************************
	2821	TCATCTCAGGGAACGTTAGCTGACAGGCAGGATGGGTACGTGTCTTCAGAAAGTGGAAGC
P AAC84422	2821	**************************************

DNA37140	2881	CACCACCAGTTTGTCACATCTTCAGGTGCTGGATTTTTCTTACCACAACATGACAGTAGT
P_AAC84422	2881	CACCACCAGTTTGTCACATCTTCAGGTGCTGGATTTTTCTTACCACAACATGACAGTAGT
DNA37140	2941	GGGACCTGCCATATTGACAATAGCAGTGAAGCTGATGTGGAAGCTGCCACAGATCTGTTC ********************************
P_AAC84422	2941	GGGACCTGCCATATTGACAATAGCAGTGAAGCTGATGTGGAAGCTGCCACAGATCTGTTC
DNA37140	3001	CTTTGTCCGTTTTTGGGATCCACAGGCCCTATGTATTTGAAGGGAAATGTGTATGGCTCA ***********************************
P_AAC84422	3001	CTTTGTCCGTTTTTGGGATCCACAGGCCCTATGTATTTGAAGGGAAATGTGTATGGCTCA
DNA37140	3061	GATCCTTTTGAAACATATCATACAGGTTGCAGTCCTGACCCAAGAACAGTTTTAATGGAC ***********************************
P_AAC84422	3061	GATCCTTTTGAAACATATCATACAGGTTGCAGTCCTGACCCAAGAACAGTTTTAATGGAC
DNA37140	3121	CACTATGAGCCCAGTTACATAAAGAAAAAGGAGTGCTACCCATGTTCTCATCCTTCAGAA **********************************
P_AAC84422	3121	CACTATGAGCCCAGTTACATAAAGAAAAAGGAGTGCTACCCATGTTCTCATCCTTCAGAA
DNA37140	3181	GAATCCTGCGAACGGAGCTTCAGTAATATATCGTGGCCTTCACATGTGAGGAAGCTACTT *********************************
P_AAC84422	3181	GAATCCTGCGAACGGAGCTTCAGTAATATATCGTGGCCTTCACATGTGAGGAAGCTACTT
DNA37140	3241	AACACTAGTTACTCTCACAATGAAGGACCTGGAATGAAAAATCTGTGTCTAAACAAGTCC ***********************************
P_AAC84422	3241	AACACTAGTTACTCTCACAATGAAGGACCTGGAATGAAAAATCTGTGTCTAAACAAGTCC
DNA37140	3301	TCTTTAGATTTTAGTGCAAATCCAGAGCCAGCGTCGGTTGCCTCGAGTAATTCTTTCATG ************************************
P_AAC84422	3301	TCTTTAGATTTTAGTGCAAATCCAGAGCCAGCGTCGGTTGCCTCGAGTAATTCTTTCATG
DNA37140	3361	GGTACCTTTGGAAAAGCTCTCAGGAGACCTCACCTAGATGCCTATTCAAGCTTTGGACAG *********************************
P_AAC84422	3361	GGTACCTTTGGAAAAGCTCTCAGGAGACCTCACCTAGATGCCTATTCAAGCTTTGGACAG
DNA37140	3421	CCATCAGATTGTCAGCCAAGAGCCTTTTATTTGAAAGCTCATTCTTCCCCAGACTTGGAC ***********************************
P_AAC84422	3421	CCATCAGATTGTCAGCCAAGAGCCTTTTATTTGAAAGCTCATTCTTCCCCAGACTTGGAC
DNA37140	3481	TCTGGGTCAGAGGAAGAAGGAAGAAAGGACAGATTTTCAGGAAGAAAATCACATTTGT *********************************
P_AAC84422	3481	TCTGGGTCAGAGGAAGAAGGAAAGGACAGATTTTCAGGAAGAAAATCACATTTGT
DNA37140	3541	ACCTTTAAACAGACTTTAGAAAACTACAGGACTCCAAATTTTCAGTCTTATGACTTGGAC
P_AAC84422	3541	ACCTTTAAACAGACTTTAGAAAACTACAGGACTCCAAATTTTCAGTCTTATGACTTGGAC
DNA37140	3601	ACATAGACTGAATGAGACCAAAGGAAAAGCTTAACATACTACCTCAAGTGAACTTTTATT *****************************
P_AAC84422	3601	ACATAGACTGAATGAGACCAAAGGAAAAGCTTAACATACTACCTCAAGTGAACTTTTATT
DNA37140	3661	TAAAAGAGAGAATCTTATGTTTTTTAAATGGAGTTATGAATTTTAAAAGGATAAAAAT **********
P_AAC84422	3661	TAAAAGAGAGAATCTTATGTTTTTTAAATGGAGTTATGAATTTTAAAAGGATAAAAAT
DNA37140	3721	GCTTTATTTATACAGATGAACCAAAATTACAAAAAGTTATGAAAATTTTTATACTGGGAA

P AAC84422	3721	**************************************
DNA37140	3781	TGATGCTCATATAAGAATACCTTTTTAAACTATTTTTTAACTTTGTTTTATGCAAAAAAG
		*************
P_AAC84422	3/81	TGATGCTCATATAAGAATACCTTTTTAAACTATTTTTTAACTTTGTTTTATGCAAAAAAG
DNA37140	3841	TATCTTACGTAAATTAATGATATAAATCATGATTATTTTATGTATTTTATAATGCCAGA *********************************
P_AAC84422	3841	TATCTTACGTAAATTAATGATATAAATCATGATTATTTTATGTATTTTATTTA
DNA37140	3901	TTTCTTTTTATGGAAAATGAGTTACTAAAGCATTTTAAATAATACCTGCCTTGTACCATT
P_AAC84422	3901	TTTCTTTTTATGGAAAATGAGTTACTAAAGCATTTTAAATAATACCTGCCTTGTACCATT
DNA37140	3961	TTTTAAATAGAAGTTACTTCATTATATTTTGCACATTATATTTAATAAAATGTGTCAATT *********************************
P_AAC84422	3961	TTTTAAATAGAAGTTACTTCATTATATTTTGCACATTATATTTAATAAAATGTGTCAATT
DNA37140	4021	TGAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAA
P_AAC84422	4021	TGAAAAAAAAAAAAAAAAAAAAAAAA
Score = 38	07 (7	man PRO326 DNA fragment #2. (3807 bp) [1 seg] 547 bits), Expect = 0.0 07/3807 (100%), at 247,1-4053,3807, Strand +/+
DNA37140	247	ATGAGCGCGCGAGCCTCCGTGCGCGCGCGCGGGGTTGGGGCTGCTGCTGTGCGCGGTG **********
P_AAX37743	1	ATGAGCGCCGAGCCTCCGTGCGCGCGCGCGGGGTTGGGGCTGCTGCTGCGCGGTG
DNA37140	307	CTGGGGCGCCGCTGGCCGGTCCGACAGCGGCGGTCGCGGGGAACTCGGGCAGCCCTCTGGG
P_AAX37743	61	$\tt CTGGGGCGCTGGCCGGTCCGACAGCGGCGGCGGGGAACTCGGGCAGCCCTCTGGGGGGGG$
DNA37140	367	GTAGCCGCCGAGCGCCCATGCCCCACTACCTGCCGCTGCCTCGGGGACCTGCTGGACTGC ***********************************
P_AAX37743	121	$\tt GTAGCCGCCGAGCGCCCATGCCCCACTACCTGCCGCTGCCTCGGGGACCTGCTGGACTGC$
DNA37140	427	AGTCGTAAGCGGCTAGCGCGTCTTCCCGAGCCACTCCCGTCCTGGGTCGCTCGGCTGGAC
P_AAX37743	181	AGTCGTAAGCGGCTAGCGCGTCTTCCCGAGCCACTCCCGTCCTGGGTCGCTCGGCTGGAC
DNA37140	487	TTAAGTCACAACAGATTATCTTTCATCAAGGCAAGTTCCATGAGCCACCTTCAAAGCCTT
P_AAX37743	241	TTAAGTCACAACAGATTATCTTTCATCAAGGCAAGTTCCATGAGCCACCTTCAAAGCCTT
DNA37140	547	CGAGAAGTGAAACTGAACAACAATGAATTGGAGACCATTCCAAATCTGGGACCAGTCTCG
P_AAX37743	301	$\tt CGAGAAGTGAACAACAATGAATTGGAGACCATTCCAAATCTGGGACCAGTCTCG$
DNA37140	607	GCAAATATTACACTTCTCTCCTTGGCTGGAAACAGGATTGTTGAAATACTCCCTGAACAT
P_AAX37743	361	GCAAATATTACACTTCTCCTTGGCTGGAAACAGGATTGTTGAAATACTCCCTGAACAT
DNA37140	667	CTGAAAGAGTTTCAGTCCCTTGAAACTTTGGACCTTAGCAGCAACAATATTTCAGAGCTC

P_AAX37743	421	CTGAAAGAGTTTCAGTCCCTTGAAACTTTGGACCTTAGCAGCAACAATATTTCAGAGCTC
DNA37140	727	CAAACTGCATTTCCAGCCCTACAGCTCAAATATCTGTATCTCAACAGCAACCGAGTCACA
P_AAX37743	481	CAAACTGCATTTCCAGCCCTACAGCTCAAATATCTGTATCTCAACAGCAACCGAGTCACA
DNA37140	787	TCAATGGAACCTGGGTATTTTGACAATTTGGCCAACACACTCCTTGTGTTAAAGCTGAAC
P_AAX37743	541	TCAATGGAACCTGGGTATTTTGACAATTTGGCCAACACTCCTTGTGTTAAAGCTGAAC
DNA37140	847	AGGAACCGAATCTCAGCTATCCCACCCAAGATGTTTAAACTGCCCCAACTGCAACATCTC
P_AAX37743	601	AGGAACCGAATCTCAGCTATCCCACCCAAGATGTTTAAACTGCCCCAACTGCAACATCTC
DNA37140	907	GAATTGAACCGAAACAAGATTAAAAATGTAGATGGACTGACATTCCAAGGCCTTGGTGCT
P_AAX37743	661	GAATTGAACCGAAACAAGATTAAAAATGTAGATGGACTGACATTCCAAGGCCTTGGTGCT
DNA37140	967	CTGAAGTCTCTGAAAATGCAAAGAAATGGAGTAACGAAACTTATGGATGG
P_AAX37743	721	CTGAAGTCTCTGAAAATGCAAAGAAATGGAGTAACGAAACTTATGGATGG
DNA37140	1027	GGGCTGAGCAACATGGAAATTTTGCAGCTGGACCATAACAACCTAACAGAGATTACCAAA
P_AAX37743	781	GGGCTGAGCAACATGGAAATTTTGCAGCTGGACCATAACAACCTAACAGAGATTACCAAA
DNA37140	1087	GGCTGGCTTTACGGCTTGCTGATGCTGCAGGAACTTCATCTCAGCCAAAATGCCATCAAC
P_AAX37743	841	GGCTGGCTTTACGGCTTGCTGATGCTGCAGGAACTTCATCTCAGCCAAAATGCCATCAAC
DNA37140	1147	AGGATCAGCCCTGATGCCTGGGAGTTCTGCCAGAAGCTCAGTGAGCTGGACCTAACTTTC
P_AAX37743	901	AGGATCAGCCCTGATGCCTGGGAGTTCTGCCAGAAGCTCAGTGAGCTGGACCTAACTTTC
DNA37140	1207	AATCACTTATCAAGGTTAGATGATTCAAGCTTCCTTGGCCTAAGCTTACTAAATACACTG
P_AAX37743	961	AATCACTTATCAAGGTTAGATGATTCAAGCTTCCTTGGCCTAAGCTTACTAAATACACTG
DNA37140	1267	CACATTGGGAACAACAGAGTCAGCTACATTGCTGATTGTGCCTTCCGGGGGCTTTCCAGT
P_AAX37743	1021	${\tt CACATTGGGAACAACAGAGTCAGCTACATTGCTGATTGTGCCTTCCGGGGGCTTTCCAGT}$
DNA37140	1327	TTAAAGACTTTGGATCTGAAGAACAATGAAATTTCCTGGACTATTGAAGACATGAATGGT
P_AAX37743	1081	TTAAAGACTTTGGATCTGAAGAACAATGAAATTTCCTGGACTATTGAAGACATGAATGGT
DNA37140	1387	GCTTTCTCTGGGCTTGACAAACTGAGGCGACTGATACTCCAAGGAAATCGGATCCGTTCT
P_AAX37743	1141	${\tt GCTTTCTCTGGGCTTGACAAACTGAGGCGACTGATACTCCAAGGAAATCGGATCCGTTCT}$
DNA37140	1447	ATTACTAAAAAAGCCTTCACTGGTTTGGATGCATTGGAGCATCTAGACCTGAGTGACAAC
P_AAX37743	1201	ATTACTAAAAAAGCCTTCACTGGTTTGGATGCATTGGAGCATCTAGACCTGAGTGACAAC
DNA37140	1507	GCAATCATGTCTTTACAAGGCAATGCATTTTCACAAATGAAGAAACTGCAACAATTGCAT
P_AAX37743	1261	GCAATCATGTCTTTACAAGGCAATGCATTTTCACAAATGAAGAAACTGCAACAATTGCAT

DNA37140	1567	TTAAATACATCAAGCCTTTTGTGCGATTGCCAGCTAAAATGGCTCCCACAGTGGGTGG
P_AAX37743	1321	TTAAATACATCAAGCCTTTTGTGCGATTGCCAGCTAAAATGGCTCCCACAGTGGGTGG
DNA37140	1627	GAAAACAACTTTCAGAGCTTTGTAAATGCCAGTTGTGCCCATCCTCAGCTGCTAAAAGGA *******************************
P_AAX37743	1381	GAAAACAACTTTCAGAGCTTTGTAAATGCCAGTTGTGCCCATCCTCAGCTGCTAAAAGGA
DNA37140	1687	AGAAGCATTTTTGCTGTTAGCCCAGATGGCTTTGTGTGTG
P_AAX37743	1441	AGAAGCATTTTTGCTGTTAGCCCAGATGGCTTTGTGTGTG
DNA37140	1747	ATCACGGTTCAGCCAGAAACACAGTCGGCAATAAAAGGTTCCAATTTGAGTTTCATCTGC **********************************
P_AAX37743	1501	ATCACGGTTCAGCCAGAAACACAGTCGGCAATAAAAGGTTCCAATTTGAGTTTCATCTGC
DNA37140	18,07	TCAGCTGCCAGCAGCAGTGATTCCCCAATGACTTTTGCTTGGAAAAAAAGACAATGAACTA
P_AAX37743	1561	TCAGCTGCCAGCAGCAGTGATTCCCCAATGACTTTTGCTTGGAAAAAAAGACAATGAACTA
DNA37140	1867	CTGCATGATGCTGAAATGGAAAATTATGCACACCTCCGGGCCCAAGGTGGCGAGGTGATG
P_AAX37743	1621	CTGCATGATGCTGAAATGGAAAATTATGCACACCTCCGGGCCCAAGGTGGCGAGGTGATG
DNA37140	1927	GAGTATACCACCATCCTTCGGCTGCGCGAGGTGGAATTTGCCAGTGAGGGGAAATATCAG
P_AAX37743	1681	GAGTATACCACCATCCTTCGGCTGCGCGAGGTGGAATTTGCCAGTGAGGGGAAATATCAG
DNA37140	1987	TGTGTCATCTCCAATCACTTTGGTTCATCCTACTCTGTCAAAGCCAAGCTTACAGTAAAT
P_AAX37743	1741	TGTGTCATCTCCAATCACTTTGGTTCATCCTACTCTGTCAAAGCCAAGCTTACAGTAAAT
DNA37140	2047	ATGCTTCCCTCATTCACCAAGACCCCCATGGATCTCACCATCCGAGCTGGGGCCATGGCA ***********************************
P_AAX37743	1801	ATGCTTCCCTCATTCACCAAGACCCCCATGGATCTCACCATCCGAGCTGGGGCCATGGCA
DNA37140	2107	CGCTTGGAGTGTGCTGTGGGGCACCCAGCCCCCAGATAGCCTGGCAGAAGGATGGG
P_AAX37743	1861	CGCTTGGAGTGTGCTGTGGGGCACCCAGCCCCCAGATAGCCTGGCAGAAGGATGGG
DNA37140	2167	GGCACAGACTTCCCAGCTGCACGGGAGAGACGCATGCATG
P_AAX37743	1921	GGCACAGACTTCCCAGCTGCACGGGAGAGACGCATGCATG
DNA37140	2227	TTCTTTATCGTGGATGTGAAGATAGAGGACATTGGGGTATACAGCTGCACAGCTCAGAAC
P_AAX37743	1981	TTCTTTATCGTGGATGTGAAGATAGAGGACATTGGGGTATACAGCTGCACAGCTCAGAAC
DNA37140	2287	AGTGCAGGAAGTATTTCAGCAAATGCAACTCTGACTGTCCTAGAAACACCATCATTTTTG ********************************
P_AAX37743	2041	AGTGCAGGAAGTATTTCAGCAAATGCAACTCTGACTGTCCTAGAAACACCATCATTTTTG
DNA37140	2347	CGGCCACTGTTGGACCGAACTGTAACCAAGGGAGAAACAGCCGTCCTACAGTGCATTGCT
P_AAX37743	2101	CGGCCACTGTTGGACCGAACTGTAACCAAGGGAGAAACAGCCGTCCTACAGTGCATTGCT

DNA37140	2407	GGAGGAAGCCCTCCCCTAAACTGAACTGGACCAAAGATGATAGCCCATTGGTGGTAACC
P_AAX37743	2161	GGAGGAAGCCCTCCCCCTAAACTGAACTGGACCAAAGATGATAGCCCATTGGTGGTAACC
DNA37140	2467	GAGAGGCACTTTTTTGCAGCAGGCAATCAGCTTCTGATTATTGTGGACTCAGATGTCAGT ************************************
P_AAX37743	2221	GAGAGGCACTTTTTTGCAGCAGGCAATCAGCTTCTGATTATTGTGGACTCAGATGTCAGT
DNA37140	2527	GATGCTGGGAAATACACATGTGAGATGTCTAACACCCTTGGCACTGAGAGAGGAAACGTG ***********************************
P_AAX37743	2281	GATGCTGGGAAATACACATGTGAGATGTCTAACACCCTTGGCACTGAGAGAGA
DNA37140	2587	CGCCTCAGTGTGATCCCCACTCCAACCTGCGACTCCCCTCAGATGACAGCCCCATCGTTA ***********************************
P_AAX37743	2341	CGCCTCAGTGTGATCCCCACTCCAACCTGCGACTCCCCTCAGATGACAGCCCCATCGTTA
DNA37140	2647	GACGATGACGGATGGGCCACTGTGGGTGTCGTGATCATAGCCGTGGTTTGCTGTGTGGTG ******************
P_AAX37743	2401	GACGATGACGGATGGGCCACTGTGGGTGTCGTGATCATAGCCGTGGTTTGCTGTGTGGTG
DNA37140	2707	GGCACGTCACTCGTGTGGGTGGTCATCATATACCACACAAGGCGGAGGAATGAAGATTGC ***********************************
P_AAX37743	2461	GGCACGTCACTCGTGTGGGTGGTCATCATATACCACACAAGGCGGAGGAATGAAGATTGC
DNA37140	2767	AGCATTACCAACACAGATGAGACCAACTTGCCAGCAGATATTCCTAGTTATTTGTCATCT
P_AAX37743	2521	AGCATTACCAACACAGATGAGACCAACTTGCCAGCAGATATTCCTAGTTATTTGTCATCT
DNA37140	2827	CAGGGAACGTTAGCTGACAGGCAGGATGGGTACGTGTCTTCAGAAAGTGGAAGCCACCAC ***************************
P_AAX37743	2581	CAGGGAACGTTAGCTGACAGGÇAGGATGGGTACGTGTCTTCAGAAAGTGGAAGCCACCAC
DNA37140	2887	CAGTTTGTCACATCTTCAGGTGCTGGATTTTTCTTACCACAACATGACAGTAGTGGGACC *********************************
P_AAX37743	2641	CAGTTTGTCACATCTTCAGGTGCTGGATTTTTCTTACCACAACATGACAGTAGTGGGACC
DNA37140	2947	TGCCATATTGACAATAGCAGTGAAGCTGATGTGGAAGCTGCCACAGATCTGTTCCTTTGT
P_AAX37743	2701	TGCCATATTGACAATAGCAGTGAAGCTGATGTGGAAGCTGCCACAGATCTGTTCCTTTGT
DNA37140	3007	CCGTTTTTGGGATCCACAGGCCCTATGTATTTGAAGGGAAATGTGTATGGCTCAGATCCT
P_AAX37743	2761	CCGTTTTTGGGATCCACAGGCCCTATGTATTTGAAGGGAAATGTGTATGGCTCAGATCCT
DNA37140	3067	TTTGAAACATATCATACAGGTTGCAGTCCTGACCCAAGAACAGTTTTAATGGACCACTAT *******************************
P_AAX37743	2821	TTTGAAACATATCATACAGGTTGCAGTCCTGACCCAAGAACAGTTTTAATGGACCACTAT
DNA37140	3127	GAGCCCAGTTACATAAAGAAAAAGGAGTGCTACCCATGTTCTCATCCTTCAGAAGAATCC **********************************
P_AAX37743	2881	GAGCCCAGTTACATAAAGAAAAAGGAGTGCTACCCATGTTCTCATCCTTCAGAAGAATCC
DNA37140	3187	TGCGAACGGAGCTTCAGTAATATCGTGGCCTTCACATGTGAGGAAGCTACTTAACACT ***************************
P_AAX37743	2941	TGCGAACGGAGCTTCAGTAATATCGTGGCCTTCACATGTGAGGAAGCTACTTAACACT
DNA37140	3247	AGTTACTCTCACAATGAAGGACCTGGAATGAAAAATCTGTGTCTAAACAAGTCCTCTTTA

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P_AAX37743	3001	AGTTACTCTCACAATGAAGGACCTGGAATGAAAAATCTGTGTCTAAACAAGTCCTCTTTA
DNA37140	3307	GATTTTAGTGCAAATCCAGAGCCAGCGTCGGTTGCCTCGAGTAATTCTTTCATGGGTACC
P_AAX37743	3061	GATTTTAGTGCAAATCCAGAGCCAGCGTCGGTTGCCTCGAGTAATTCTTTCATGGGTACC
DNA37140	3367	TTTGGAAAAGCTCTCAGGAGACCTCACCTAGATGCCTATTCAAGCTTTGGACAGCCATCA
P_AAX37743	3121	TTTGGAAAAGCTCTCAGGAGACCTCACCTAGATGCCTATTCAAGCTTTGGACAGCCATCA
DNA37140	3427	GATTGTCAGCCAAGAGCCTTTTATTTGAAAGCTCATTCTTCCCCAGACTTGGACTCTGGG
P_AAX37743	3181	GATTGTCAGCCAAGAGCCTTTTATTTGAAAGCTCATTCTTCCCCAGACTTGGACTCTGGG
DNA37140	3487	TCAGAGGAAGATGGGAAAGAAAGGACAGATTTTCAGGAAGAAAATCACATTTGTACCTTT
P_AAX37743	3241	TCAGAGGAAGAAGGAAAGAAAGGACAGATTTTCAGGAAGAAAATCACATTTGTACCTTT
DNA37140	3547	AAACAGACTTTAGAAAACTACAGGACTCCAAATTTTCAGTCTTATGACTTGGACACATAG
P_AAX37743	3301	AAACAGACTTTAGAAAACTACAGGACTCCAAATTTTCAGTCTTATGACTTGGACACATAG
DNA37140	3607	ACTGAATGAGACCAAAGGAAAAGCTTAACATACTACCTCAAGTGAACTTTTATTTA
P_AAX37743	3361	ACTGAATGAGACCAAAGGAAAAGCTTAACATACTACCTCAAGTGAACTTTTATTTA
DNA37140	3667	AGAGAGAATCTTATGTTTTTTAAATGGAGTTATGAATTTTAAAAGGATAAAAATGCTTTA
P_AAX37743	3421	AGAGAGAATCTTATGTTTTTTAAATGGAGTTATGAATTTTAAAAGGATAAAAATGCTTTA
DNA37140	3727	TTTATACAGATGAACCAAAATTACAAAAAGTTATGAAAATTTTTATACTGGGAATGATGC
P_AAX37743	3481	TTTATACAGATGAACCAAAATTACAAAAAGTTATGAAAATTTTTATACTGGGAATGATGC
DNA37140	3787	TCATATAAGAATACCTTTTTAAACTATTTTTTAACTTTTTTTT
P_AAX37743	3541	TCATATAAGAATACCTTTTTAAACTATTTTTTAACTTTTGTTTTATGCAAAAAAGTATCTT
DNA37140	3847	ACGTAAATTAATGATATAAATCATGATTATTTTATGTATTTTTATAATGCCAGATTTCTT
P_AAX37743	3601	ACGTAAATTAATGATATAAATCATGATTATTTTATGTATTTTTATAATGCCAGATTTCTT
DNA37140	3907	TTTATGGAAAATGAGTTACTAAAGCATTTTAAATAATACCTGCCTTGTACCATTTTTTAA
P_AAX37743	3661	TTTATGGAAAATGAGTTACTAAAGCATTTTAAATAATACCTGCCTTGTACCATTTTTTAA
DNA37140	3967	ATAGAAGTTACTTCATTATATTTTGCACATTATATTTAATAAAATGTGTCAATTTGAAAA
P_AAX37743	3721	ATAGAAGTTACTTCATTATATTTTGCACATTATATTTAATAAAATGTGTCAATTTGAAAA
DNA37140	4027	AAAAAAAAAAAAAAAAAAAAA ****************
P_AAX37743	3781	ААААААААААААААААААААА
>8 P_AAF7242	22 Hum	nan PRO335 cDNA. (3662 bp) [1 seg]

<sup>&</sup>gt;8 P\_AAF72422 Human PRO335 cDNA. (3662 bp) [1 seg] Score = 3542 (7022 bits), Expect = 0.0

Identities	= 35	42/3542 (100%), at 483,121-4024,3662, Strand +/+
DNA37140	483	GGACTTAAGTCACAACAGATTATCTTTCATCAAGGCAAGTTCCATGAGCCACCTTCAAAG
P_AAF72422	121	GGACTTAAGTCACAACAGATTATCTTTCATCAAGGCAAGTTCCATGAGCCACCTTCAAAG
DNA37140	543	CCTTCGAGAAGTGAAACTGAACAACAATGAATTGGAGACCATTCCAAATCTGGGACCAGT
P_AAF72422	181	CCTTCGAGAAGTGAAACTGAACAACAATGAATTGGAGACCATTCCAAATCTGGGACCAGT
DNA37140	603	CTCGGCAAATATTACACTTCTCTCTTGGCTGGAAACAGGATTGTTGAAATACTCCCTGA
P_AAF72422	241	CTCGGCAAATATTACACTTCTCCTTGGCTGGAAACAGGATTGTTGAAATACTCCCTGA
DNA37140	663	ACATCTGAAAGAGTTTCAGTCCCTTGAAACTTTGGACCTTAGCAGCAACAATATTTCAGA
P_AAF72422	301	ACATCTGAAAGAGTTTCAGTCCCTTGAAACTTTGGACCTTAGCAGCAACAATATTTCAGA
DNA37140	723	GCTCCAAACTGCATTTCCAGCCCTACAGCTCAAATATCTGTATCTCAACAGCAACCGAGT
P_AAF72422	361	GCTCCAAACTGCATTTCCAGCCCTACAGCTCAAATATCTGTATCTCAACAGCAACCGAGT
DNA37140	783	CACATCAATGGAACCTGGGTATTTTGACAATTTGGCCAACACACTCCTTGTGTTAAAGCT
P_AAF72422	421	CACATCAATGGAACCTGGGTATTTTGACAATTTGGCCAACACACTCCTTGTGTTAAAGCT
DNA37140	843	GAACAGGAACCGAATCTCAGCTATCCCACCCAAGATGTTTAAACTGCCCCAACTGCAACA
P_AAF72422	481	GAACAGGAACCGAATCTCAGCTATCCCACCCAAGATGTTTAAACTGCCCCAACTGCAACA
DNA37140	903	TCTCGAATTGAACCGAAACAAGATTAAAAATGTAGATGGACTGACATTCCAAGGCCTTGG
P_AAF72422	541	TCTCGAATTGAACCGAAACAAGATTAAAAATGTAGATGGACTGACATTCCAAGGCCTTGG
DNA37140	963	TGCTCTGAAGTCTCTGAAAATGCAAAGAAATGGAGTAACGAAACTTATGGATGG
P_AAF72422	601	TGCTCTGAAGTCTCTGAAAATGCAAAGAAATGGAGTAACGAAACTTATGGATGG
DNA37140	1023	TTGGGGGCTGAGCAACATGGAAATTTTGCAGCTGGACCATAACAACCTAACAGAGATTAC
P_AAF72422	661	TTGGGGGCTGAGCAACATGGAAATTTTGCAGCTGGACCATAACAACCTAACAGAGATTAC
DNA37140	1083	CAAAGGCTGGCTTTACGGCTTGCTGATGCTGCAGGAACTTCATCTCAGCCAAAATGCCAT
P_AAF72422	721	CAAAGGCTGGCTTTACGGCTTGCTGATGCTGCAGGAACTTCATCTCAGCCAAAATGCCAT
DNA37140	1143	CAACAGGATCAGCCCTGATGCCTGGGAGTTCTGCCAGAAGCTCAGTGAGCTGGACCTAAC
P_AAF72422	781	CAACAGGATCAGCCCTGATGCCTGGGAGTTCTGCCAGAAGCTCAGTGAGCTGGACCTAAC
DNA37140	1203	TTTCAATCACTTATCAAGGTTAGATGATTCAAGCTTCCTTGGCCTAAGCTTACTAAATAC
P_AAF72422	841	TTTCAATCACTTATCAAGGTTAGATGATTCAAGCTTCCTTGGCCTAAGCTTACTAAATAC
DNA37140	1263	ACTGCACATTGGGAACAACAGAGTCAGCTACATTGCTGATTGTGCCTTCCGGGGGCTTTC
P_AAF72422	901	ACTGCACATTGGGAACAACAGAGTCAGCTACATTGCTGATTGTGCCTTCCGGGGGCTTTC

DNA37140	1323	CAGTTTAAAGACTTTGGATCTGAAGAACAATGAAATTTCCTGGACTATTGAAGACATGAA
P_AAF72422	961	CAGTTTAAAGACTTTGGATCTGAAGAACAATGAAATTTCCTGGACTATTGAAGACATGAA
DNA37140	1383	TGGTGCTTTCTCTGGGCTTGACAAACTGAGGCGACTGATACTCCAAGGAAATCGGATCCG ***********************************
P_AAF72422	1021	TGGTGCTTTCTCTGGGCTTGACAAACTGAGGCGACTGATACTCCAAGGAAATCGGATCCG
DNA37140	1443	TTCTATTACTAAAAAAGCCTTCACTGGTTTGGATGCATTGGAGCATCTAGACCTGAGTGA ********************************
P_AAF72422	1081	TTCTATTACTAAAAAAGCCTTCACTGGTTTGGATGCATTGGAGCATCTAGACCTGAGTGA
DNA37140	1503	CAACGCAATCATGTCTTTACAAGGCAATGCATTTTCACAAATGAAGAAACTGCAACAATT *******************************
P_AAF72422	1141	CAACGCAATCATGTCTTTACAAGGCAATGCATTTTCACAAATGAAGAAACTGCAACAATT
DNA37140	1563	GCATTTAAATACATCAAGCCTTTTGTGCGATTGCCAGCTAAAATGGCTCCCACAGTGGGT ********************************
P_AAF72422	1201	GCATTTAAATACATCAAGCCTTTTGTGCGATTGCCAGCTAAAATGGCTCCCACAGTGGGT
DNA37140	1623	GGCGGAAACAACTTTCAGAGCTTTGTAAATGCCAGTTGTGCCCATCCTCAGCTGCTAAA **********************************
P_AAF72422	1261	GGCGGAAAACAACTTTCAGAGCTTTGTAAATGCCAGTTGTGCCCATCCTCAGCTGCTAAA
DNA37140	1683	AGGAAGAAGCATTTTTGCTGTTAGCCCAGATGGCTTTGTGTGTG
P_AAF72422	1321	AGGAAGAAGCATTTTTGCTGTTAGCCCAGATGGCTTTGTGTGTG
DNA37140	1743	CCAGATCACGGTTCAGCCAGAAACACAGTCGGCAATAAAAGGTTCCAATTTGAGTTTCAT
P_AAF72422	1381	CCAGATCACGGTTCAGCCAGAAACACAGTCGGCAATAAAAGGTTCCAATTTGAGTTTCAT
DNA37140	1803	CTGCTCAGCTGCCAGCAGCAGTGATTCCCCAATGACTTTTGCTTGGAAAAAAGACAATGA **********************************
P_AAF72422	1441	CTGCTCAGCTGCCAGCAGCAGTGATTCCCCCAATGACTTTTGCTTGGAAAAAAAGACAATGA
DNA37140	1863	ACTACTGCATGATGCTGAAATGGAAAATTATGCACACCTCCGGGCCCAAGGTGGCGAGGT
P_AAF72422	1501	ACTACTGCATGATGCTGAAATGGAAAATTATGCACACCTCCGGGCCCAAGGTGGCGAGGT
DNA37140	1923	GATGGAGTATACCACCATCCTTCGGCTGCGCGAGGTGGAATTTGCCAGTGAGGGGAAATA *************************
P_AAF72422	1561	GATGGAGTATACCACCATCCTTCGGCTGCGCGAGGTGGAATTTGCCAGTGAGGGGAAATA
DNA37140	1983	TCAGTGTGTCATCTCCAATCACTTTGGTTCATCCTACTCTGTCAAAGCCAAGCTTACAGT ************************************
P_AAF72422	1621	TCAGTGTGTCATCTCCAATCACTTTGGTTCATCCTACTCTGTCAAAGCCAAGCTTACAGT
DNA37140	2043	AAATATGCTTCCCTCATTCACCAAGACCCCCATGGATCTCACCATCCGAGCTGGGGCCAT ***********************************
P_AAF72422	1681	AAATATGCTTCCCTCATTCACCAAGACCCCCATGGATCTCACCATCCGAGCTGGGGCCAT
DNA37140	2103	GGCACGCTTGGAGTGTGCTGCTGTGGGGCACCCAGCCCCCAGATAGCCTGGCAGAAGGA *****************************
P_AAF72422	1741	GGCACGCTTGGAGTGTGCTGCTGTGGGGCACCCAGCCCCCAGATAGCCTGGCAGAAGGA

DNA37140	2163	TGGGGGCACAGACTTCCCAGCTGCACGGGAGAGACGCATGCAT
P_AAF72422	1801	TGGGGGCACAGACTTCCCAGCTGCACGGGAGAGACGCATGCAT
DNA37140	2223	CGTGTTCTTTATCGTGGATGTGAAGATAGAGGACATTGGGGTATACAGCTGCACAGCTCA
P_AAF72422	1861	$\tt CGTGTTCTTTATCGTGGATGTGAAGATAGAGGACATTGGGGTATACAGCTGCACAGCTCA$
DNA37140	2283	GAACAGTGCAGGAAGTATTTCAGCAAATGCAACTCTGACTGTCCTAGAAACACCATCATT
P_AAF72422	1921	GAACAGTGCAGGAAGTATTTCAGCAAATGCAACTCTGACTGTCCTAGAAACACCATCATT
DNA37140	2343	TTTGCGGCCACTGTTGGACCGAACTGTAACCAAGGGAGAAACAGCCGTCCTACAGTGCAT ************************************
P_AAF72422	1981	TTTGCGGCCACTGTTGGACCGAACTGTAACCAAGGGAGAAACAGCCGTCCTACAGTGCAT
DNA37140	2403	TGCTGGAGGAAGCCCTCCCCCTAAACTGAACTGGACCAAAGATGATAGCCCATTGGTGGT
P_AAF72422	2041	TGCTGGAGGAAGCCCTCCCCCTAAACTGAACTGGACCAAAGATGATAGCCCATTGGTGGT
DNA37140	2463	AACCGAGAGCACTTTTTTGCAGCAGGCAATCAGCTTCTGATTATTGTGGACTCAGATGT *********************************
P_AAF72422	2101	AACCGAGAGGCACTTTTTTGCAGCAGGCAATCAGCTTCTGATTATTGTGGACTCAGATGT
DNA37140	2523	CAGTGATGCTGGGAAATACACATGTGAGATGTCTAACACCCTTGGCACTGAGAGGAAA
P_AAF72422	2161	CAGTGATGCTGGGAAATACACATGTGAGATGTCTAACACCCTTGGCACTGAGAGAGA
DNA37140	2583	CGTGCGCCTCAGTGTGATCCCCACTCCAACCTGCGACTCCCCTCAGATGACAGCCCCATC
P_AAF72422	2221	CGTGCGCCTCAGTGTGATCCCÇACTCCAACCTGCGACTCCCCTCAGATGACAGCCCCATC
DNA37140	2643	GTTAGACGATGACGGATGGGCCACTGTGGGTGTCGTGATCATAGCCGTGGTTTGCTGTGT
P_AAF72422	2281	GTTAGACGATGACGGATGGGCCACTGTGGGTGTCGTGATCATAGCCGTGGTTTGCTGTGT
DNA37140	2703	GGTGGGCACGTCACTCGTGTGGGTGGTCATCATATACCACACAAGGCGGAGGAATGAAGA **************************
P_AAF72422	2341	GGTGGGCACGTCACTCGTGTGGGTGGTCATCATATACCACACAAGGCGGAGGAATGAAGA
DNA37140	2763	TTGCAGCATTACCAACACAGATGAGACCAACTTGCCAGCAGATATTCCTAGTTATTTGTC *********************************
P_AAF72422	2401	TTGCAGCATTACCAACACAGATGAGACCAACTTGCCAGCAGATATTCCTAGTTATTTGTC
DNA37140	2823	ATCTCAGGGAACGTTAGCTGACAGGCAGGATGGGTACGTGTCTTCAGAAAGTGGAAGCCA *******************************
P_AAF72422	2461	ATCTCAGGGAACGTTAGCTGACAGGCAGGATGGGTACGTGTCTTCAGAAAGTGGAAGCCA
DNA37140	2883	CCACCAGTTTGTCACATCTTCAGGTGCTGGATTTTTCTTACCACAACATGACAGTAGTGG
P_AAF72422	2521	CCACCAGTTTGTCACATCTTCAGGTGCTGGATTTTTCTTACCACAACATGACAGTAGTGG
DNA37140	2943	GACCTGCCATATTGACAATAGCAGTGAAGCTGATGTGGAAGCTGCCACAGATCTGTTCCT ******************************
P_AAF72422	2581	GACCTGCCATATTGACAATAGCAGTGAAGCTGATGTGGAAGCTGCCACAGATCTGTTCCT
DNA37140	3003	TTGTCCGTTTTTGGGATCCACAGGCCCTATGTATTTGAAGGGAAATGTGTATGGCTCAGA

P_AAF72422	2641	TTGTCCGTTTTTGGGATCCACAGGCCCTATGTATTTGAAGGGAAATGTGTATGGCTCAGA
DNA37140	3063	TCCTTTTGAAACATATCATACAGGTTGCAGTCCTGACCCAAGAACAGTTTTAATGGACCA
P_AAF72422	2701	TCCTTTTGAAACATATCATACAGGTTGCAGTCCTGACCCAAGAACAGTTTTAATGGACCA
DNA37140	3123	CTATGAGCCCAGTTACATAAAGAAAAAGGAGTGCTACCCATGTTCTCATCCTTCAGAAGA
P_AAF72422	2761	CTATGAGCCCAGTTACATAAAGAAAAAGGAGTGCTACCCATGTTCTCATCCTTCAGAAGA
DNA37140	3183	ATCCTGCGAACGGAGCTTCAGTAATATATCGTGGCCTTCACATGTGAGGAAGCTACTTAA
P_AAF72422	2821	ATCCTGCGAACGGAGCTTCAGTAATATCGTGGCCTTCACATGTGAGGAAGCTACTTAA
DNA37140	3243	CACTAGTTACTCTCACAATGAAGGACCTGGAATGAAAAATCTGTGTCTAAACAAGTCCTC
P_AAF72422	2881	CACTAGTTACTCTCACAATGAAGGACCTGGAATGAAAAATCTGTGTCTAAACAAGTCCTC
DNA37140	3303	TTTAGATTTTAGTGCAAATCCAGAGCCAGCGTCGGTTGCCTCGAGTAATTCTTTCATGGG
P_AAF72422	2941	TTTAGATTTTAGTGCAAATCCAGAGCCAGCGTCGGTTGCCTCGAGTAATTCTTTCATGGG
DNA37140	3363	TACCTTTGGAAAAGCTCTCAGGAGACCTCACCTAGATGCCTATTCAAGCTTTGGACAGCC
P_AAF72422	3001	TACCTTTGGAAAAGCTCTCAGGAGACCTCACCTAGATGCCTATTCAAGCTTTGGACAGCC
DNA37140	3423	ATCAGATTGTCAGCCAAGAGCCTTTTATTTGAAAGCTCATTCTTCCCCAGACTTGGACTC
P_AAF72422	3061	ATCAGATTGTCAGCCAAGAGCCTTTTATTTGAAAGCTCATTCTTCCCCAGACTTGGACTC
DNA37140	3483	TGGGTCAGAGGAAGATGGGAAAAGGAAGGACAGATTTTCAGGAAGAAAATCACATTTGTAC
P_AAF72422	3121	TGGGTCAGAGGAAGAAGGAAGAAAGGACAGATTTTCAGGAAGAAAATCACATTTGTAC
DNA37140	3543	CTTTAAACAGACTTTAGAAAACTACAGGACTCCAAATTTTCAGTCTTATGACTTGGACAC
P_AAF72422	3181	CTTTAAACAGACTTTAGAAAACTACAGGACTCCAAATTTTCAGTCTTATGACTTGGACAC
DNA37140	3603	ATAGACTGAATGAGACCAAAGGAAAAGCTTAACATACTACCTCAAGTGAACTTTTATTTA
P_AAF72422	3241	$\verb  ************************************$
DNA37140	3663	AAAGAGAGAGAATCTTATGTTTTTTAAATGGAGTTATGAATTTTAAAAGGATAAAAATGC
P_AAF72422	3301	**************************************
DNA37140	3723	TTTATTTATACAGATGAACCAAAATTACAAAAAGTTATGAAAATTTTTATACTGGGAATG
P_AAF72422	3361	${\tt ************************************$
DNA37140	3783	ATGCTCATATAAGAATACCTTTTTAAACTATTTTTTAACTTTGTTTTATGCAAAAAAGTA
P_AAF72422	3421	**************************************
DNA37140	3843	TCTTACGTAAATTAATGATATAAATCATGATTATTTTATGTATTTTTATAATGCCAGATT
		*****



## /home/ruby/va/Molbio/carpenda/temp1/ss.DNA37140 (4053 bp)

1 P_AAF72424 2 P_AAZ52208 3 P_AAX52266 4 AX098387 5 AX056647 6 P_AAC84422 7 P_AAX37743 8 P_AAF72422 9 P_AAZ52206 10 P_AAX52264	Acing High-scoring Segment Pairs: Frame Score Match Pct E-val Human PRO326 cDNA. + 4053 4053 100 0.0 Human PRO326 cDNA clone DNA37140-1234. + 4053 4053 100 0.0 Sequence 13 from Patent W00119991. + 4053 4053 100 0.0 Sequence 3 from Patent W00075316. + 4053 4053 100 0.0 Human PRO326 polypeptide encoding cDNA. + 4021 4045 100 0.0 Human PRO326 DNA fragment #2. + 3807 3807 100 0.0 Human PRO335 cDNA. + 3542 3542 100 0.0 Protein PRO335 cDNA clone DNA41388-1234. + 3542 3542 100 0.0 Human PRO335 DNA fragment #2. + 3542 3542 100 0.0 Sequence 9 from Patent W00119991. + 3542 3542 100 0.0
Score = $4053$	Numan PRO326 cDNA. (4053 bp) [1 seg] 8034 bits), Expect = 0.0 053/4053 (100%), at 1,1-4053,4053, Strand +/+
DNA37140	1 AGCCGACGCTGCTCAAGCTGCAACTCTGTTGCAGTTGGCAGTTCTTTTCGGTTTCCCTGT
P_AAF72424	1 AGCCGACGCTGCTCAAGCTGCAACTCTGTTGCAGTTGGCAGTTCTTTTCGGTTTCCCTCC
DNA37140 6	1 TGCTGTTTGGGGGCATGAAAGGGCTTCGCCGCCGGGAGTAAAAGAAGAAGGAATTGACCG
P_AAF72424 6	1 TGCTGTTTGGGGGCATGAAAGGGCTTCGCCGCCGGGAGTAAAAGAAGGAATTGACCGCGC
DNA37140 12	1 AGCGCGAGGAGGAGCGCGCACGCGACCGCGAGGGCGGGC
P_AAF72424 12	1 AGCGCGAGGAGGGCGCGCACGCGAGGGCGGGCGTGCACCCTCGGCTGGAAGT
DNA37140 18	1 TTGTGCCGGGCCCCGAGCGCGCCGGCTGGGAGCTTCGGGTAGAGACCTAGGCCGCTGG
P_AAF72424 18	1 TTGTGCCGGGCCCGAGCGCGCCGGCTGGGAGCTTCGGGTAGAGACCTAGGCCGCTGG
DNA37140 24	1 ACCGCGATGAGCGCCGAGCCTCCGTGCGCGCGCGCGGGGTTGGGGCTGCTGTGC
P_AAF72424 24	1 ACCGCGATGAGCGCGCGAGCCTCCGTGCGCGCGCGCGGGGGTTGGGGGCTGCTGTGC
DNA37140 30	1 GCGGTGCTGGGCGCTGGCCGGTCCGACAGCGGCGGTCGCGGGGAACTCGGGCAGCCC
P_AAF72424 30	**************************************
DNA37140 36	1 TCTGGGGTAGCCGCCGAGCGCCCATGCCCCACTACCTGCCGCTGCCTCGGGGACCTGCTG
P_AAF72424 36	**************************************
DNA37140 42	1 GACTGCAGTCGTAAGCGGCTAGCGCGTCTTCCCGAGCCACTCCCGTCCTGGGTCGCTCGG
P_AAF72424 42	**************************************
DNA37140 48	1 CTGGACTTAAGTCACAACAGATTATCTTTCATCAAGGCAAGTTCCATGAGCCACCTTCAA
P_AAF72424 48	**************************************

DNA37140	541	AGCCTTCGAGAAGTGAAACTGAACAACAATGAATTGGAGACCATTCCAAATCTGGGACCA ********************************
P_AAF72424	541	${\tt AGCCTTCGAGAAGTGAACTGAACAATGAATTGGAGACCATTCCAAATCTGGGACCA} \\ {\tt -}$
DNA37140	601	GTCTCGGCAAATATTACACTTCTCTCTTGGCTGGAAACAGGATTGTTGAAATACTCCCT
P_AAF72424	601	GTCTCGGCAAATATTACACTTCTCTCTTGGCTGGAAACAGGATTGTTGAAATACTCCCT
DNA37140	661	GAACATCTGAAAGAGTTTCAGTCCCTTGAAACTTTGGACCTTAGCAGCAACAATATTTCA
P_AAF72424	661	GAACATCTGAAAGAGTTTCAGTCCCTTGAAACTTTGGACCTTAGCAGCAACAATATTTCA
DNA37140	721	GAGCTCCAAACTGCATTTCCAGCCCTACAGCTCAAATATCTGTATCTCAACAGCAACCGA
P_AAF72424	721	GAGCTCCAAACTGCATTTCCAGCCCTACAGCTCAAATATCTGTATCTCAACAGCAACCGA
DNA37140	781	GTCACATCAATGGAACCTGGGTATTTTGACAATTTGGCCAACACACTCCTTGTGTTAAAG
P_AAF72424	781	GTCACATCAATGGAACCTGGGTATTTTGACAATTTGGCCAACACACTCCTTGTGTTAAAG
DNA37140	841	CTGAACAGGAACCGAATCTCAGCTATCCCACCCAAGATGTTTAAACTGCCCCAACTGCAA
P_AAF72424	841	CTGAACAGGAACCGAATCTCAGCTATCCCACCCAAGATGTTTAAACTGCCCCAACTGCAA
DNA37140	901	CATCTCGAATTGAACCGAAACAAGATTAAAAATGTAGATGGACTGACATTCCAAGGCCTT
P_AAF72424	901	CATCTCGAATTGAACCGAAACAAGATTAAAAATGTAGATGGACTGACATTCCAAGGCCTT
DNA37140	961	GGTGCTCTGAAGTCTCTGAAAATGCAAAGAAATGGAGTAACGAAACTTATGGATGG
P_AAF72424	961	GGTGCTCTGAAGTCTCTGAAAATGCAAAGAAATGGAGTAACGAAACTTATGGATGG
DNA37140	1021	TTTTGGGGGCTGAGCAACATGGAAATTTTGCAGCTGGACCATAACAACCTAACAGAGATT
P_AAF72424	1021	TTTTGGGGGCTGAGCAACATGGAAATTTTGCAGCTGGACCATAACAACCTAACAGAGA
DNA37140	1081	ACCAAAGGCTGGCTTTACGGCTTGCTGATGCTGCAGGAACTTCATCTCAGCCAAAATGCC
P_AAF72424	1081	ACCAAAGGCTGGCTTTACGGCTTGCTGATGCTGCAGGAACTTCATCTCAGCCAAAATGCC
DNA37140	1141	ATCAACAGGATCAGCCCTGATGCCTGGGAGTTCTGCCAGAAGCTCAGTGAGCTGGACCTA **********************************
P_AAF72424	1141	ATCAACAGGATCAGCCCTGATGCCTGGGAGTTCTGCCAGAAGCTCAGTGAGCTGGACCTA
DNA37140	1201	ACTTTCAATCACTTATCAAGGTTAGATGATTCAAGCTTCCTTGGCCTAAGCTTACTAAAT
P_AAF72424	1201	ACTTTCAATCACTTATCAAGGTTAGATGATTCAAGCTTCCTTGGCCTAAGCTTACTAAAT
DNA37140	1261	ACACTGCACATTGGGAACAACAGAGTCAGCTACATTGCTGATTGTGCCTTCCGGGGGCTT
P_AAF72424	1261	ACACTGCACATTGGGAACAACAGAGTCAGCTACATTGCTGATTGTGCCTTCCGGGGGCCTT
DNA37140	1321	TCCAGTTTAAAGACTTTGGATCTGAAGAACAATGAAATTTCCTGGACTATTGAAGACATG
P_AAF72424	1321	TCCAGTTTAAAGACTTTGGATCTGAAGAACAATGAAATTTCCTGGACTATTGAAGACATG
DNA37140	1381	AATGGTGCTTTCTCTGGGCTTGACAAACTGAGGCGACTGATACTCCAAGGAAATCGGATC

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P_AAF72424	1381	AATGGTGCTTTCTCTGGGCTTGACAAACTGAGGCGACTGATACTCCAAGGAAATCGGATC
DNA37140	1441	CGTTCTATTACTAAAAAGCCTTCACTGGTTTGGATGCATTGGAGCATCTAGACCTGAGT
P_AAF72424	1441	CGTTCTATTACTAAAAAAGCCTTCACTGGTTTGGATGCATTGGAGCATCTAGACCTGAGT
DNA37140	1501	GACAACGCAATCATGTCTTTACAAGGCAATGCATTTTCACAAATGAAGAAACTGCAACAA ********************************
P_AAF72424	1501	GACAACGCAATCATGTCTTTACAAGGCAATGCATTTTCACAAATGAAGAAACTGCAACAA
DNA37140	1561	TTGCATTTAAATACATCAAGCCTTTTGTGCGATTGCCAGCTAAAATGGCTCCCACAGTGG **********************************
P_AAF72424	1561	TTGCATTTAAATACATCAAGCCTTTTGTGCGATTGCCAGCTAAAATGGCTCCCACAGTGG
DNA37140	1621	GTGGCGGAAAACAACTTTCAGAGCTTTGTAAATGCCAGTTGTGCCCATCCTCAGCTGCTA ************************************
P_AAF72424	1621	GTGGCGGAAAACAACTTTCAGAGCTTTGTAAATGCCAGTTGTGCCCATCCTCAGCTGCTA
DNA37140	1681	AAAGGAAGAAGCATTTTTGCTGTTAGCCCAGATGGCTTTGTGTGTG
P_AAF72424	1681	${\tt AAAGGAAGCATTTTTGCTGTTAGCCCAGATGGCTTTGTGTGTG$
DNA37140	1741	CCCCAGATCACGGTTCAGCCAGAAACACAGTCGGCAATAAAAGGTTCCAATTTGAGTTTC
P_AAF72424	1741	CCCCAGATCACGGTTCAGCCAGAAACACAGTCGGCAATAAAAGGTTCCAATTTGAGTTTC
DNA37140	1801	ATCTGCTCAGCTGCCAGCAGCAGTGATTCCCCAATGACTTTTGCTTGGAAAAAAGACAAT **********************************
P_AAF72424	1801	ATCTGCTCAGCTGCCAGCAGCAGTGATTCCCCAATGACTTTTGCTTGGAAAAAAAGACAAT
DNA37140	1861	GAACTACTGCATGATGCTGAAATTGGAAAATTATGCACACCTCCGGGCCCAAGGTGGCGAG *********************************
P_AAF72424	1861	GAACTACTGCATGATGCTGAAATGGAAAATTATGCACACCTCCGGGCCCAAGGTGGCGAG
DNA37140	1921	GTGATGGAGTATACCACCATCCTTCGGCTGCGCGAGGTGGAATTTGCCAGTGAGGGGAAA ***************************
P_AAF72424	1921	GTGATGGAGTATACCACCATCCTTCGGCTGCGCGAGGTGGAATTTGCCAGTGAGGGGAAA
DNA37140	1981	TATCAGTGTGTCATCTCCAATCACTTTGGTTCATCCTACTCTGTCAAAGCCAAGCTTACA *********************************
P_AAF72424	1981	TATCAGTGTGTCATCTCCAATCACTTTGGTTCATCCTACTCTGTCAAAGCCAAGCTTACA
DNA37140	2041	GTAAATATGCTTCCCTCATTCACCAAGACCCCCATGGATCTCACCATCCGAGCTGGGGCC ******************************
P_AAF72424	2041	GTAAATATGCTTCCCTCATTCACCAAGACCCCCATGGATCTCACCATCCGAGCTGGGGCC
DNA37140	2101	ATGGCACGCTTGGAGTGTGCTGCTGTGGGGCACCCAGCCCCCAGATAGCCTGGCAGAAG *******************************
P_AAF72424	2101	ATGGCACGCTTGGAGTGTGCTGCTGTGGGGCACCCAGCCCCCAGATAGCCTGGCAGAAG
DNA37140	2161	GATGGGGGCACAGACTTCCCAGCTGCACGGGAGAGACGCATGCAT
P_AAF72424	2161	GATGGGGGCACAGACTTCCCAGCTGCACGGGAGAGACGCATGCAT
DNA37140	2221	GACGTGTTCTTTATCGTGGATGTGAAGATAGAGGACATTGGGGTATACAGCTGCACAGCT ************************************

P_AAF72424	2221	GACGTGTTCTTTATCGTGGATGTGAAGATAGAGGACATTGGGGTATACAGCTGCACAGCT
DNA37140	2281	CAGAACAGTGCAGGAAGTATTTCAGCAAATGCAACTCTGACTGTCCTAGAAACACCATCA
P_AAF72424	2281	CAGAACAGTGCAGGAAGTATTTCAGCAAATGCAACTCTGACTGTCCTAGAAACACCATCA
DNA37140	2341	TTTTTGCGGCCACTGTTGGACCGAACTGTAACCAAGGGAGAAACAGCCGTCCTACAGTGC ***********************************
P_AAF72424	2341	TTTTTGCGGCCACTGTTGGACCGAACTGTAACCAAGGGAGAAACAGCCGTCCTACAGTGC
DNA37140	2401	ATTGCTGGAGGAAGCCCTCCCCCTAAACTGAACTGGACCAAAGATGATAGCCCATTGGTG
P_AAF72424	2401	${\tt ATTGCTGGAGGAAGCCCTCCCCCTAAACTGAACTGGACCAAAGATGATAGCCCATTGGTG} {\tt } \\$
DNA37140	2461	GTAACCGAGAGGCACTTTTTTGCAGCAGGCAATCAGCTTCTGATTATTGTGGACTCAGA
P_AAF72424	2461	GTAACCGAGAGGCACTTTTTTGCAGCAGGCAATCAGCTTCTGATTATTGTGGACTCAGA
DNA37140	2521	GTCAGTGATGCTGGGAAATACACATGTGAGATGTCTAACACCCTTGGCACTGAGAGAGGCA
P_AAF72424	2521	GTCAGTGATGCTGGGAAATACACATGTGAGATGTCTAACACCCTTGGCACTGAGAGAGA
DNA37140	2581	AACGTGCGCCTCAGTGTGATCCCCACTCCAACCTGCGACTCCCCTCAGATGACAGCCC
P_AAF72424	2581	AACGTGCGCCTCAGTGTGATCCCCACTCCAACCTGCGACTCCCCTCAGATGACAGCCCCA
DNA37140	2641	TCGTTAGACGATGACGGATGGGCCACTGTGGGTGTCGTGATCATAGCCGTGGTTTGCTGT ************************
P_AAF72424	2641	TCGTTAGACGATGACGGATGGGCCACTGTGGGTGTCGTGATCATAGCCGTGGTTTGCTGT
DNA37140	2701	GTGGTGGGCACGTCACTCGTGTGGGTGGTCATCATATACCACACAAGGCGGAGGAATGAA
P_AAF72424	2701	GTGGTGGGCACGTCACTCGTGTGGGTGGTCATCATATACCACACAAGGCGGAGGAATGAA
DNA37140	2761	GATTGCAGCATTACCAACACAGATGAGACCAACTTGCCAGCAGATATTCCTAGTTATTTG *****************************
P_AAF72424	2761	GATTGCAGCATTACCAACACAGATGAGACCAACTTGCCAGCAGATATTCCTAGTTATTTG
DNA37140	2821	TCATCTCAGGGAACGTTAGCTGACAGGCAGGATGGGTACGTGTCTTCAGAAAGTGGAAGC *********************************
P_AAF72424	2821	TCATCTCAGGGAACGTTAGCTGACAGGCAGGATGGGTACGTGTCTTCAGAAAGTGGAAGC
DNA37140	2881	CACCACCAGTTTGTCACATCTTCAGGTGCTGGATTTTTCTTACCACAACATGACAGTAGT
P_AAF72424	2881	CACCACCAGTTTGTCACATCTTCAGGTGCTGGATTTTTCTTACCACAACATGACAGTAGT
DNA37140	2941	GGGACCTGCCATATTGACAATAGCAGTGAAGCTGATGTGGAAGCTGCCACAGATCTGTTC
P_AAF72424	2941	GGGACCTGCCATATTGACAATAGCAGTGAAGCTGATGTGGAAGCTGCCACAGATCTGTTC
DNA37140	3001	CTTTGTCCGTTTTTGGGATCCACAGGCCCTATGTATTTGAAGGGAAATGTGTATGGCTCA
P_AAF72424	3001	CTTTGTCCGTTTTTGGGATCCACAGGCCCTATGTATTTGAAGGGAAATGTGTATGGCTCA
DNA37140	3061	GATCCTTTTGAAACATATCATACAGGTTGCAGTCCTGACCCAAGAACAGTTTTAATGGAC
P AAF72424	3061	GATCCTTTTGAAACATATCATACAGGTTGCAGTCCTGACCCAAGAACAGTTTTAATGGAC

DNA37140	3121	CACTATGAGCCCAGTTACATAAAGAAAAAGGAGTGCTACCCATGTTCTCATCCTTCAGAA
P_AAF72424	3121	CACTATGAGCCCAGTTACATAAAGAAAAAGGAGTGCTACCCATGTTCTCATCCTTCAGAA
DNA37140	3181	GAATCCTGCGAACGGAGCTTCAGTAATATATCGTGGCCTTCACATGTGAGGAAGCTACTT
P_AAF72424	3181	GAATCCTGCGAACGGAGCTTCAGTAATATATCGTGGCCTTCACATGTGAGGAAGCTACTT
DNA37140	3241	AACACTAGTTACTCTCACAATGAAGGACCTGGAATGAAAAATCTGTGTCTAAACAAGTCC ***********************************
P_AAF72424	3241	AACACTAGTTACTCTCACAATGAAGGACCTGGAATGAAAAATCTGTGTCTAAACAAGTCC
DNA37140	3301	TCTTTAGATTTTAGTGCAAATCCAGAGCCAGCGTCGGTTGCCTCGAGTAATTCTTTCATG
P_AAF72424	3301	${\tt TCTTTAGATTTTAGTGCAAATCCAGAGCCAGCGTCGGTTGCCTCGAGTAATTCTTTCATG}$
DNA37140	3361	GGTACCTTTGGAAAAGCTCTCAGGAGACCTCACCTAGATGCCTATTCAAGCTTTGGACAG
P_AAF72424	3361	GGTACCTTTGGAAAAGCTCTCAGGAGACCTCACCTAGATGCCTATTCAAGCTTTGGACAG
DNA37140	3421	CCATCAGATTGTCAGCCAAGAGCCTTTTATTTGAAAGCTCATTCTTCCCCAGACTTGGAC
P_AAF72424	3421	${\tt CCATCAGATTGTCAGCCAAGAGCCTTTTATTTGAAAGCTCATTCTTCCCCAGACTTGGAC}$
DNA37140	3481	TCTGGGTCAGAGGAAGAAGGAAGAAAGGACAGATTTTCAGGAAGAAAATCACATTTGT
P_AAF72424	3481	TCTGGGTCAGAGGAAGAAGGAAGAAAGGACAGATTTTCAGGAAGAAAATCACATTTGT
DNA37140	3541	ACCTTTAAACAGACTTTAGAAAACTACAGGACTCCAAATTTTCAGTCTTATGACTTGGAC
P_AAF72424	3541	ACCTTTAAACAGACTTTAGAAAACTACAGGACTCCAAATTTTCAGTCTTATGACTTGGAC
DNA37140	3601	ACATAGACTGAATGAGACCAAAGGAAAAGCTTAACATACTACCTCAAGTGAACTTTTATT
P_AAF72424	3601	ACATAGACTGAATGAGACCAAAGGAAAAGCTTAACATACTACCTCAAGTGAACTTTTATT
DNA37140	3661	TAAAAGAGAGAGAATCTTATGTTTTTTAAATGGAGTTATGAATTTTAAAAGGATAAAAAT
P_AAF72424	3661	TAAAAGAGAGAGAATCTTATGTTTTTTAAATGGAGTTATGAATTTTAAAAGGATAAAAAT
DNA37140	3721	GCTTTATTTATACAGATGAACCAAAATTACAAAAAGTTATGAAAATTTTTATACTGGGAA
P_AAF72424	3721	GCTTTATTTATACAGATGAACCAAAATTACAAAAAGTTATGAAAATTTTTATACTGGGAA
DNA37140	3781	TGATGCTCATATAAGAATACCTTTTTAAACTATTTTTTTAACTTTGTTTTATGCAAAAAAG
P_AAF72424	3781	TGATGCTCATATAAGAATACCTTTTTAAACTATTTTTTAACTTTGTTTTATGCAAAAAAG
DNA37140	3841	TATCTTACGTAAATTAATGATATAAATCATGATTATTTTATGTATTTTTATAATGCCAGA
P_AAF72424	3841	TATCTTACGTAAATTAATGATATAAATCATGATTATTTTATGTATTTTATAATGCCAGA
DNA37140	3901	TTTCTTTTTATGGAAAATGAGTTACTAAAGCATTTTAAATAATACCTGCCTTGTACCATT
P_AAF72424	3901	TTTCTTTTTATGGAAATGAGTTACTAAAGCATTTTAAATAATACCTGCCTTGTACCATT

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3961 TTTTAAATAGAAGTTACTTCATTATATTTTGCACATTATATTTAATAAAATGTGTCAATT
  DNA37140
               *******************
          3961 TTTTAAATAGAAGTTACTTCATTATATTTTGCACATTATATTTAATAAAATGTGTCAATT
 P AAF72424
  DNA37140
          4021 TGAAAAAAAAAAAAAAAAAAAAAAAAAAA
P AAF72424
          4021 TGAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAA
>2 P AAZ52208 Human PRO326 protein encoding cDNA, UNQ287. DNA, PAT 18-JUL-2000
 (4053 bp) [1 seg]
 Score = 4053 (8034 bits), Expect = 0.0
 Identities = 4053/4053 (100%), at 1,1-4053,4053, Strand +/+
  DNA37140
             1 \ \ \mathsf{AGCCGACGCTGCTCAAGCTGCAACTCTGTTGCAGTTGGCAGTTCTTTTCGGTTTCCCTCC}
               ********************
P AAZ52208
             1 AGCCGACGCTGCTCAAGCTGCAACTCTGTTGCAGTTGGCAGTTCTTTTCGGTTTCCCTCC
            61 TGCTGTTTGGGGGCATGAAAGGGCTTCGCCGCCGGGAGTAAAAGAAGGAATTGACCGGGC
  DNA37140
P AAZ52208 ·
            61 TGCTGTTTGGGGGCATGAAAGGGCTTCGCCGCCGGGAGTAAAAGAAGGAATTGACCGGGC
  DNA37140
           121 AGCGCGAGGGAGGCGCGCACGCGACGCGAGGGCGGGCGTGCACCCTCGGCTGGAAGT
              ***********************
P AAZ52208
           121 AGCGCGAGGGAGGAGCGCGCACGCGACCGCGAGGGCGGGCGTGCACCCTCGGCTGGAAGT
           181 TTGTGCCGGGCCCCGAGCGCGCCGGCTGGGAGCTTCGGGTAGAGACCTAGGCCGCTGG
  DNA37140
              **********
P AAZ52208
           181 TTGTGCCGGGCCCGAGCGCCGCCGGCTGGAGCTTCGGGTAGAGACCTAGGCCGCTGG
  DNA37140
           241 ACCGCGATGAGCGCCGAGCCTCCGTGCGCGCGCGCGGGGTTGGGGCTGCTGTGC
              *****************
P AAZ52208
           241 ACCGCGATGAGCGCCGAGCCTCCGTGCGCGCGCGCGGGGTTGGGGCTGCTGTGC
  DNA37140
           301 GCGGTGCTGGGGGGCGCTGGCCGGTCCGACAGCGGGGGGAACTCGGGCAGCCC
              ******************
P AAZ52208
           301 GCGGTGCTGGGCGCCTGGCCGGTCCGACAGCGGCGGTCGCGGGGAACTCGGGCAGCCC
  DNA37140
           361 TCTGGGGTAGCCGCCGAGCGCCCATGCCCACTACCTGCCGCTGCCTCGGGGACCTGCTG
              ******************
P AAZ52208
           361 TCTGGGGTAGCCGCCGAGCGCCCATGCCCACTACCTGCCGCTGCCTCGGGGACCTGCTG
 DNA37140
           421 GACTGCAGTCGTAAGCGGCTAGCGCGTCTTCCCGAGCCACTCCCGTCCTGGGTCGCTCGG
              ******************
P AAZ52208
           421 GACTGCAGTCGTAAGCGGCTAGCGCGTCTTCCCGAGCCACTCCCGTCCTGGGTCGCTCGG
 DNA37140
          481 CTGGACTTAAGTCACAACAGATTATCTTTCATCAAGGCAAGTTCCATGAGCCACCTTCAA
              ************
P AAZ52208
          481 CTGGACTTAAGTCACAACAGATTATCTTTCATCAAGGCAAGTTCCATGAGCCACCTTCAA
 DNA37140
          541 AGCCTTCGAGAAGTGAAACTGAACAATGAATTGGAGACCATTCCAAATCTGGGACCA
              **************
P AAZ52208
          541 AGCCTTCGAGAAGTGAAACTGAACAACAATGAATTGGAGACCATTCCAAATCTGGGACCA
 DNA37140
          601 GTCTCGGCAAATATTACACTTCTCTCTTGGCTGGAAACAGGATTGTTGAAATACTCCCT
              P AAZ52208
          601 GTCTCGGCAAATATTACACTTCTCTCTTGGCTGGAAACAGGATTGTTGAAATACTCCCT
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DNA37140	661	GAACATCTGAAAGAGTTTCAGTCCCTTGAAACTTTGGACCTTAGCAGCAACAATATTTCA **************************
P_AAZ52208	661	GAACATCTGAAAGAGTTTCAGTCCCTTGAAACTTTTGGACCTTAGCAGCAACAATATTTCA
DNA37140	721	GAGCTCCAAACTGCATTTCCAGCCCTACAGCTCAAATATCTGTATCTCAACAGCAACCGA
P_AAZ52208	721	GAGCTCCAAACTGCATTTCCAGCCCTACAGCTCAAATATCTGTATCTCAACAGCAACCGA
DNA37140	781	GTCACATCAATGGAACCTGGGTATTTTGACAATTTGGCCAACACACTCCTTGTGTTAAAG
P_AAZ52208	781	GTCACATCAATGGAACCTGGGTATTTTGACAATTTGGCCAACACACTCCTTGTGTTAAAG
DNA37140	841	CTGAACAGGAACCGAATCTCAGCTATCCCACCCAAGATGTTTAAACTGCCCCAACTGCAA
P_AAZ52208	841	CTGAACAGGAACCGAATCTCAGCTATCCCACCCAAGATGTTTAAACTGCCCCAACTGCAA
DNA37140	901	CATCTCGAATTGAACCGAAACAAGATTAAAAATGTAGATGGACTGACATTCCAAGGCCTT
P_AAZ52208	901	CATCTCGAATTGAACCGAAACAAGATTAAAAATGTAGATGGACTGACATTCCAAGGCCTT
DNA37140	961	GGTGCTCTGAAGTCTCTGAAAATGCAAAGAAATGGAGTAACGAAACTTATGGATGG
P_AAZ52208	961	GGTGCTCTGAAGTCTCTGAAAATGCAAAGAAATGGAGTAACGAAACTTATGGATGG
DNA37140	1021	TTTTGGGGGCTGAGCAACATGGAAATTTTGCAGCTGGACCATAACAACCTAACAGAGATT
P_AAZ52208	1021	TTTTGGGGGCTGAGCAACATGGAAATTTTGCAGCTGGACCATAACAACCTAACAGAGATT
DNA37140	1081	ACCAAAGGCTGGCTTTACGGCTTGCTGATGCTGCAGGAACTTCATCTCAGCCAAAATGCC
P_AAZ52208	1081	${\tt ACCAAAGGCTGGCTTTACGGCTTGCTGATGCTGCAGGAACTTCATCTCAGCCAAAATGCC}$
DNA37140	1141	ATCAACAGGATCAGCCCTGATGCCTGGGAGTTCTGCCAGAAGCTCAGTGAGCTGGACCTA **********************************
P_AAZ52208	1141	ATCAACAGGATCAGCCCTGATGCCTGGGAGTTCTGCCAGAAGCTCAGTGAGCTGGACCTA
DNA37140	1201	ACTTTCAATCACTTATCAAGGTTAGATGATTCAAGCTTCCTTGGCCTAAGCTTACTAAAT
P_AAZ52208	1201	ACTTTCAATCACTTATCAAGGTTAGATGATTCAAGCTTCCTTGGCCTAAGCTTACTAAAT
DNA37140	1261	ACACTGCACATTGGGAACAACAGAGTCAGCTACATTGCTGATTGTGCCTTCCGGGGGCTT
P_AAZ52208	1261	ACACTGCACATTGGGAACAACAGAGTCAGCTACATTGCTGATTGTGCCTTCCGGGGGCTT
DNA37140	1321	TCCAGTTTAAAGACTTTGGATCTGAAGAACAATGAAATTTCCTGGACTATTGAAGACATG
P_AAZ52208	1321	TCCAGTTTAAAGACTTTGGATCTGAAGAACAATGAAATTTCCTGGACTATTGAAGACATG
DNA37140	1381	AATGGTGCTTTCTCTGGGCTTGACAAACTGAGGCGACTGATACTCCAAGGAAATCGGATC ************************************
P_AAZ52208	1381	AATGGTGCTTTCTCTGGGCTTGACAAACTGAGGCGACTGATACTCCAAGGAAATCGGATC
DNA37140	1441	CGTTCTATTACTAAAAAAGCCTTCACTGGTTTGGATGCATTGGAGCATCTAGACCTGAGT **********************************
P_AAZ52208	1441	CGTTCTATTACTAAAAAAGCCTTCACTGGTTTGGATGCATTGGAGCATCTAGACCTGAGT
DNA37140	1501	GACAACGCAATCATGTCTTTACAAGGCAATGCATTTTCACAAATGAAGAAACTGCAACAA

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P_AAZ52208	1501	GACAACGCAATCATGTCTTTACAAGGCAATGCATTTTCACAAATGAAGAAACTGCAACAA
DNA37140	1561	TTGCATTTAAATACATCAAGCCTTTTGTGCGATTGCCAGCTAAAATGGCTCCCACAGTĞG ***********************************
P_AAZ52208	1561	TTGCATTTAAATACATCAAGCCTTTTGTGCGATTGCCAGCTAAAATGGCTCCCACAGTGG
DNA37140	1621	GTGGCGGAAAACAACTTTCAGAGCTTTGTAAATGCCAGTTGTGCCCATCCTCAGCTGCTA
P_AAZ52208	1621	GTGGCGGAAAACAACTTTCAGAGCTTTGTAAATGCCAGTTGTGCCCATCCTCAGCTGCTA
DNA37140	1681	AAAGGAAGAAGCATTTTTGCTGTTAGCCCAGATGGCTTTGTGTGTG
P_AAZ52208	1681	AAAGGAAGAAGCATTTTTGCTGTTAGCCCAGATGGCTTTGTGTGTG
DNA37140	1741	CCCCAGATCACGGTTCAGCCAGAAACACAGTCGGCAATAAAAGGTTCCAATTTGAGTTTC
P_AAZ52208	1741	$\tt CCCCAGATCACGGTTCAGCCAGAAACACAGTCGGCAATAAAAGGTTCCAATTTGAGTTTC$
DNA37140	1801	ATCTGCTCAGCTGCCAGCAGCAGTGATTCCCCAATGACTTTTGCTTGGAAAAAAGACAAT
P_AAZ52208	1801	ATCTGCTCAGCTGCCAGCAGCAGTGATTCCCCCAATGACTTTTGCTTGGAAAAAAAGACAAT
DNA37140	1861	GAACTACTGCATGATGCTGAAATGGAAAATTATGCACACCTCCGGGCCCAAGGTGGCGAG
P_AAZ52208	1861	GAACTACTGCATGATGCTGAAATGGAAAATTATGCACACCTCCGGGCCCAAGGTGGCGAG
DNA37140	1921	GTGATGGAGTATACCACCATCCTTCGGCTGCGCGAGGTGGAATTTGCCAGTGAGGGGAAA
P_AAZ52208	1921	GTGATGGAGTATACCACCATCCTTCGGCTGCGCGAGGTGGAATTTGCCAGTGAGGGGAAA
DNA37140	1981	TATCAGTGTCATCTCCAATCACTTTGGTTCATCCTACTCTGTCAAAGCCAAGCTTACA
P_AAZ52208	1981	TATCAGTGTCATCTCCAATCACTTTGGTTCATCCTACTCTGTCAAAGCCAAGCTTACA
DNA37140	2041	GTAAATATGCTTCCCTCATTCACCAAGACCCCCATGGATCTCACCATCCGAGCTGGGGCC ******************************
P_AAZ52208	2041	GTAAATATGCTTCCCTCATTCACCAAGACCCCCATGGATCTCACCATCCGAGCTGGGGCC
DNA37140	2101	ATGGCACGCTTGGAGTGTGCTGCTGTGGGGCACCCAGCCCCCCAGATAGCCTGGCAGAAG
P_AAZ52208	2101	ATGGCACGCTTGGAGTGTGCTGCTGTGGGGCACCCAGCCCCCCAGATAGCCTGGCAGAAG
DNA37140	2161	GATGGGGGCACAGACTTCCCAGCTGCACGGGAGAGACGCATGCAT
P_AAZ52208	2161	GATGGGGGCACAGACTTCCCAGCTGCACGGGAGAGACGCATGCAT
DNA37140	2221	GACGTGTTCTTTATCGTGGATGTGAAGATAGAGGACATTGGGGTATACAGCTGCACAGCT
P_AAZ52208	2221	GACGTGTTCTTTATCGTGGATGTGAAGATAGAGGACATTGGGGTATACAGCTGCACAGCT
DNA37140	2281	CAGAACAGTGCAGGAAGTATTTCAGCAAATGCAACTCTGACTGTCCTAGAAACACCATCA
P_AAZ52208	2281	CAGAACAGTGCAGGAAGTATTTCAGCAAATGCAACTCTGACTGTCCTAGAAACACCATCA
DNA37140	2341	TTTTTGCGGCCACTGTTGGACCGAACTGTAACCAAGGGAGAAACAGCCGTCCTACAGTGC

P_AAZ52208	2341	TTTTTGCGGCCACTGTTGGACCGAACTGTAACCAAGGGAGAAACAGCCGTCCTACAGTGC
DNA37140	2401	ATTGCTGGAGGAAGCCCTCCCCCTAAACTGAACTGGACCAAAGATGATAGCCCATTGGTG
P_AAZ52208	2401	ATTGCTGGAGGAAGCCCTCCCCCTAAACTGAACTGGACCAAAGATGATAGCCCATTGGTG
DNA37140	2461	GTAACCGAGAGGCACTTTTTTGCAGCAGGCAATCAGCTTCTGATTATTGTGGACTCAGAT
P_AAZ52208	2461	GTAACCGAGAGGCACTTTTTTGCAGCAGGCAATCAGCTTCTGATTATTGTGGACTCAGAT
DNA37140	2521	GTCAGTGATGCTGGGAAATACACATGTGAGATGTCTAACACCCTTGGCACTGAGAGAGA
P_AAZ52208	2521	GTCAGTGATGCTGGGAAATACACATGTGAGATGTCTAACACCCTTGGCACTGAGAGAGGA
DNA37140	2581	AACGTGCGCCTCAGTGTGATCCCCACTCCAACCTGCGACTCCCCTCAGATGACAGCCCCA *****************************
P_AAZ52208	2581	AACGTGCGCCTCAGTGTGATCCCCACTCCAACCTGCGACTCCCCTCAGATGACAGCCCCA
DNA37140	2641	TCGTTAGACGATGACGGATGGGCCACTGTGGGTGTCGTGATCATAGCCGTGGTTTGCTGT ************************
P_AAZ52208	2641	TCGTTAGACGATGACGGATGGGCCACTGTGGGTGTCGTGATCATAGCCGTGGTTTGCTGT
DNA37140	2701	GTGGTGGCACGTCACTCGTGTGGGTGGTCATCATATACCACACAAGGCGGAGGAATGAA ****************************
P_AAZ52208	2701	GTGGTGGCACGTCACTCGTGTGGGTGGTCATCATATACCACACAAGGCGGAGGAATGAA
DNA37140	2761	GATTGCAGCATTACCAACACAGATGAGACCAACTTGCCAGCAGATATTCCTAGTTATTTG *****************************
P_AAZ52208	2761	GATTGCAGCATTACCAACACAGATGAGACCAACTTGCCAGCAGATATTCCTAGTTATTTG
DNA37140	2821	TCATCTCAGGGAACGTTAGCTGACAGGCAGGATGGGTACGTGTCTTCAGAAAGTGGAAGC *********************************
P_AAZ52208	2821	TCATCTCAGGGAACGTTAGCTGACAGGCAGGATGGGTACGTGTCTTCAGAAAGTGGAAGC
DNA37140	2881	CACCACCAGTTTGTCACATCTTCAGGTGCTGGATTTTTCTTACCACAACATGACAGTAGT *********************************
P_AAZ52208	2881	CACCACCAGTTTGTCACATCTTCAGGTGCTGGATTTTTCTTACCACAACATGACAGTAGT
DNA37140	2941	GGGACCTGCCATATTGACAATAGCAGTGAAGCTGATGTGGAAGCTGCCACAGATCTGTTC ********************************
P_AAZ52208	2941	GGGACCTGCCATATTGACAATAGCAGTGAAGCTGATGTGGAAGCTGCCACAGATCTGTTC
DNA37140		CTTTGTCCGTTTTTGGGATCCACAGGCCCTATGTATTTGAAGGGAAATGTGTATGGCTCA ***********************************
P_AAZ52208		CTTTGTCCGTTTTTGGGATCCACAGGCCCTATGTATTTGAAGGGAAATGTGTATGGCTCA
DNA37140		GATCCTTTTGAAACATATCATACAGGTTGCAGTCCTGACCCAAGAACAGTTTTAATGGAC ***********************************
P_AAZ52208		GATCCTTTTGAAACATATCATACAGGTTGCAGTCCTGACCCAAGAACAGTTTTAATGGAC
DNA37140		CACTATGAGCCCAGTTACATAAAGAAAAAGGAGTGCTACCCATGTTCTCATCCTTCAGAA **********************************
P_AAZ52208		CACTATGAGCCCAGTTACATAAAGAAAAAGGAGTGCTACCCATGTTCTCATCCTTCAGAA
DNA37140		GAATCCTGCGAACGGAGCTTCAGTAATATCGTGGCCTTCACATGTGAGGAAGCTACTT *********************************
P_AAZ52208	3181	GAATCCTGCGAACGGAGCTTCAGTAATATATCGTGGCCTTCACATGTGAGGAAGCTACTT

DNA37140	3241	AACACTAGTTACTCTCACAATGAAGGACCTGGAATGAAAAATCTGTGTCTAAACAAGTCC
P_AAZ52208	3241	AACACTAGTTACTCTCACAATGAAGGACCTGGAATGAAAAATCTGTGTCTAAACAAGTCC
DNA37140	3301	TCTTTAGATTTTAGTGCAAATCCAGAGCCAGCGTCGGTTGCCTCGAGTAATTCTTTCATG
P_AAZ52208	3301	
DNA37140	3361	GGTACCTTTGGAAAAGCTCTCAGGAGACCTCACCTAGATGCCTATTCAAGCTTTGGACAG
P_AAZ52208	3361	GGTACCTTTGGAAAAGCTCTCAGGAGACCTCACCTAGATGCCTATTCAAGCTTTGGACAG
DNA37140	3421	CCATCAGATTGTCAGCCAAGAGCCTTTTATTTGAAAGCTCATTCTTCCCCAGACTTGGAC
P_AAZ52208	3421	CCATCAGATTGTCAGCCAAGAGCCTTTTATTTGAAAGCTCATTCTTCCCCAGACTTGGAC
DNA37140	3481	TCTGGGTCAGAGGAAGATGGGAAAGAAGGACAGATTTTCAGGAAGAAAATCACATTTGT
P_AAZ52208	3481	TCTGGGTCAĞAGAAGAAGGAAGAAAGGACAGATTTTCAGGAAGAAAATCACATTTGT
DNA37140	3541	ACCTTTAAACAGACTTTAGAAAACTACAGGACTCCAAATTTTCAGTCTTATGACTTGGAC
P_AAZ52208	3541	ACCTTTAAACAGACTTTAGAAAACTACAGGACTCCAAATTTTCAGTCTTATGACTTGGAC
DNA37140	3601	ACATAGACTGAATGAGACCAAAGGAAAAGCTTAACATACTACCTCAAGTGAACTTTTATT
P_AAZ52208	3601	ACATAGACTGAATGAGACCAAAGGAAAAGCTTAACATACTACCTCAAGTGAACTTTTATT
DNA37140	3661	TAAAAGAGAGAGAATCTTATGTTTTTTAAATGGAGTTATGAATTTTAAAAGGATAAAAAT
P_AAZ52208	3661	${\tt ************************************$
DNA37140	3721	GCTTTATTTATACAGATGAACCAAAATTACAAAAAGTTATGAAAATTTTTATACTGGGAA
P_AAZ52208	3721	GCTTTATTTATACAGATGAACCAAAATTACAAAAAGTTATGAAAATTTTTATACTGGGAA
DNA37140	3781	TGATGCTCATATAAGAATACCTTTTTAAACTATTTTTTAACTTTTTTATGCAAAAAAG
P_AAZ52208	3781	TGATGCTCATATAAGAATACCTTTTTTAAACTATTTTTTAACTTTTTTATGCAAAAAAG
DNA37140	3841	TATCTTACGTAAATTAATGATATAAATCATGATTATTTTATGTATTTTTATAATGCCAGA
P_AAZ52208	3841	TATCTTACGTAAATTAATGATATAAATCATGATTATTTTATGTATTTTTATAATGCCAGA
DNA37140	3901	TTTCTTTTTATGGAAAATGAGTTACTAAAGCATTTTAAATAATACCTGCCTTGTACCATT
P_AAZ52208	3901	TTTCTTTTTATGGAAAATGAGTTACTAAAGCATTTTAAATAATACCTGCCTTGTACCATT
DNA37140	3961	TTTTAAATAGAAGTTACTTCATTATATTTTGCACATTATATTTAATAAAATGTGTCAATT
P_AAZ52208	3961	TTTTAAATAGAAGTTACTTCATTATATTTTGCACATTATATTTAATAAAATGTGTCAATT
DNA37140	4021	TGAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAA
P_AAZ52208	4021	TGAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAA

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>3 P AAX52266 Protein PRO326 cDNA clone DNA37140-1234. DNA, PAT 25-JUN-1999
(405\overline{3} \text{ bp}) [1 \text{ seg}]
 Score = 4053 (8034 bits), Expect = 0.0
 Identities = 4053/4053 (100%), at 1,1-4053,4053, Strand +/+
  DNA37140
            1 AGCCGACGCTGCTCAAGCTGCAACTCTGTTGCAGTTGGCAGTTCTTTTCGGTTTCCCTCC
              *******************
            1 AGCCGACGCTGCTCAAGCTGCAACTCTGTTGCAGTTGGCAGTTCTTTTCGGTTTCCCTCC
P AAX52266
  DNA37140
           61 TGCTGTTTGGGGGCATGAAAGGGCTTCGCCGCCGGGAGTAAAAGAAGGAATTGACCGGGC
              *******************
           61 TGCTGTTTGGGGGCATGAAAGGGCTTCGCCGCCGGGAGTAAAAGAAGAATTGACCGGGC
P AAX52266
  DNA37140
          ********************
P AAX52266
          121 AGCGCGAGGGAGGGCGCGCGCGACCGCGAGGGCGGGCGTGCACCCTCGGCTGGAAGT
 DNA37140
          181 TTGTGCCGGGCCCGAGCGCGCCGGCTGGGAGCTTCGGGTAGAGACCTAGGCCGCTGG
P AAX52266
          181 TTGTGCCGGGCCCCGAGCGCGCCGGCTGGGAGCTTCGGGTAGAGACCTAGGCCGCTGG
 DNA37140
          241 ACCGCGATGAGCGCCGAGCCTCCGTGCGCGCGCGCGGGGTTGGGGCTGCTGTGC
             ***********************
P AAX52266
          241 ACCGCGATGAGCGCCGAGCCTCCGTGCGCGCGCGGGGGTTGGGGCTGCTGTGC
          DNA37140
             **********
P AAX52266
          301 GCGGTGCTGGGCGCGCTGGCCGGTCCGACAGCGGCGGTCGCGGGGAACTCGGGCAGCCC
 DNA37140
          361 TCTGGGGTAGCCGCCGAGCGCCCATGCCCCACTACCTGCCGCTGCCTCGGGGACCTGCTG
             *****************
P AAX52266
          361 TCTGGGGTAGCCGCCGAGCGCCCATGCCCACTACCTGCCGCTGCCTCGGGGACCTGCTG
          421 GACTGCAGTCGTAAGCGGCTAGCGCGTCTTCCCGAGCCACTCCCGTCCTGGGTCGCTCGG
 DNA37140
             *****************
P AAX52266
          421 GACTGCAGTCGTAAGCGGCTAGCGCGTCTTCCCGAGCCACTCCCGTCCTGGGTCGCTCGG
          481 CTGGACTTAAGTCACAACAGATTATCTTTCATCAAGGCAAGTTCCATGAGCCACCTTCAA
 DNA37140
             P AAX52266
          481 CTGGACTTAAGTCACAACAGATTATCTTTCATCAAGGCAAGTTCCATGAGCCACCTTCAA
 DNA37140
          541 AGCCTTCGAGAAGTGAAACTGAACAACAATGAATTGGAGACCATTCCAAATCTGGGACCA
             *******************
P AAX52266
          541 AGCCTTCGAGAAGTGAAACTGAACAACAATGAATTGGAGACCATTCCAAATCTGGGACCA
 DNA37140
          601 GTCTCGGCAAATATTACACTTCTCTCTTGGCTGGAAACAGGATTGTTGAAATACTCCCT
             *******************
P AAX52266
          601 GTCTCGGCAAATATTACACTTCTCTCTTGGCTGGAAACAGGATTGTTGAAATACTCCCT
 DNA37140
          661 GAACATCTGAAAGAGTTTCAGTCCCTTGAAACTTTGGACCTTAGCAGCAACAATATTTCA
             *******
P AAX52266
          661 GAACATCTGAAAGAGTTTCAGTCCCTTGAAACTTTGGACCTTAGCAGCAACAATATTTCA
 DNA37140
          721 GAGCTCCAAACTGCATTTCCAGCCCTACAGCTCAAATATCTGTATCTCAACAGCAACCGA
             P AAX52266
          721 GAGCTCCAAACTGCATTTCCAGCCCTACAGCTCAAATATCTGTATCTCAACAGCAACCGA
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DNA37140	781	GTCACATCAATGGAACCTGGGTATTTTGACAATTTGGCCAACACACTCCTTGTGTTAAAG
P_AAX52266	781	GTCACATCAATGGAACCTGGGTATTTTGACAATTTGGCCAACACACTCCTTGTGTTAAAG
DNA37140	841	CTGAACAGGAACCGAATCTCAGCTATCCCACCCAAGATGTTTAAACTGCCCCAACTGCAA
P_AAX52266	841	CTGAACAGGAACCGAATCTCAGCTATCCCACCCAAGATGTTTAAACTGCCCCAACTGCAA
DNA37140	901	CATCTCGAATTGAACCGAAACAAGATTAAAAATGTAGATGGACTGACATTCCAAGGCCTT
P_AAX52266	901	CATCTCGAATTGAACCGAAACAAGATTAAAAATGTAGATGGACTGACATTCCAAGGCCTT
DNA37140	961	GGTGCTCTGAAGTCTCTGAAAATGCAAAGAAATGGAGTAACGAAACTTATGGATGG
P_AAX52266	961	GGTGCTCTGAAGTCTCTGAAAATGCAAAGAAATGGAGTAACGAAACTTATGGATGG
DNA37140	1021	TTTTGGGGGCTGAGCAACATGGAAATTTTGCAGCTGGACCATAACAACCTAACAGAGATT ********************************
P_AAX52266	1021	TTTTGGGGGCTGAGCAACATGGAAATTTTGCAGCTGGACCATAACAACCTAACAGAGATT
DNA37140	1081	ACCAAAGGCTGGCTTTACGGCTTGCTGATGCTGCAGGAACTTCATCTCAGCCAAAATGCC ***********************************
P_AAX52266	1081	ACCAAAGGCTGGCTTTACGGCTTGCTGATGCTGCAGGAACTTCATCTCAGCCAAAATGCC
DNA37140	1141	ATCAACAGGATCAGCCCTGATGCCTGGGAGTTCTGCCAGAAGCTCAGTGAGCTGGACCTA **********************************
P_AAX52266	1141	ATCAACAGGATCAGCCCTGATGCCTGGGAGTTCTGCCAGAAGCTCAGTGAGCTGGACCTA
DNA37140	1201	ACTTTCAATCACTTATCAAGGTTAGATGATTCAAGCTTCCTTGGCCTAAGCTTACTAAAT
P_AAX52266	1201	ACTTTCAATCACTTATCAAGGTTAGATGATTCAAGCTTCCTTGGCCTAAGCTTACTAAAT
DNA37140	1261	ACACTGCACATTGGGAACAACAGAGTCAGCTACATTGCTGATTGTGCCTTCCGGGGGCTT ********************
P_AAX52266	1261	ACACTGCACATTGGGAACAACAGAGTCAGCTACATTGCTGATTGTGCCTTCCGGGGGCTT
DNA37140	1321	TCCAGTTTAAAGACTTTGGATCTGAAGAACAATGAAATTTCCTGGACTATTGAAGACATG
P_AAX52266	1321	TCCAGTTTAAAGACTTTGGATCTGAAGAACAATGAAATTTCCTGGACTATTGAAGACATG
DNA37140	1381	AATGGTGCTTTCTCTGGGCTTGACAAACTGAGGCGACTGATACTCCAAGGAAATCGGATC ************************************
P_AAX52266	1381	AATGGTGCTTTCTCTGGGCTTGACAAACTGAGGCGACTGATACTCCAAGGAAATCGGATC
DNA37140	1441	CGTTCTATTACTAAAAAAGCCTTCACTGGTTTGGATGCATTGGAGCATCTAGACCTGAGT **********************************
P_AAX52266	1441	CGTTCTATTACTAAAAAAGCCTTCACTGGTTTGGATGCATTGGAGCATCTAGACCTGAGT
DNA37140	1501	GACAACGCAATCATGTCTTTACAAGGCAATGCATTTTCACAAATGAAGAAACTGCAACAA
P_AAX52266	1501	GACAACGCAATCATGTCTTTACAAGGCAATGCATTTTCACAAATGAAGAAACTGCAACAA
DNA37140	1561	TTGCATTTAAATACATCAAGCCTTTTGTGCGATTGCCAGCTAAAATGGCTCCCACAGTGG **********************************
P_AAX52266	1561	TTGCATTTAAATACATCAAGCCTTTTGTGCGATTGCCAGCTAAAATGGCTCCCACAGTGG
DNA37140	1621	GTGGCGGAAAACAACTTTCAGAGCTTTGTAAATGCCAGTTGTGCCCATCCTCAGCTGCTA

P_AAX52266	1621	GTGGCGGAAAACAACTTTCAGAGCTTTGTAAATGCCAGTTGTGCCCATCCTCAGCTGCT
DNA37140	1681	AAAGGAAGAAGCATTTTTGCTGTTAGCCCAGATGGCTTTGTGTGTG
P_AAX52266	1681	AAAGGAAGAAGCATTTTTGCTGTTAGCCCAGATGGCTTTGTGTGTG
DNA37140	1741	CCCCAGATCACGGTTCAGCCAGAAACACAGTCGGCAATAAAAGGTTCCAATTTGAGTTTC
P_AAX52266	1741	CCCCAGATCACGGTTCAGCCAGAAACACAGTCGGCAATAAAAGGTTCCAATTTGAGTTTC
DNA37140	1801	ATCTGCTCAGCTGCCAGCAGCAGTGATTCCCCAATGACTTTTGCTTGGAAAAAAGACAAT
P_AAX52266	1801	ATCTGCTCAGCTGCCAGCAGCAGTGATTCCCCCAATGACTŢTTGCTTGGAAAAAAGACAAT
DNA37140	1861	GAACTACTGCATGATGCTGAAATGGAAAATTATGCACACCTCCGGGCCCAAGGTGGCGAG
P_AAX52266	1861	GAACTACTGCATGATGCTGAAATGGAAAATTATGCACACCTCCGGGCCCAAGGTGGCGAG
DNA37140	1921	GTGATGGAGTATACCACCATCCTTCGGCTGCGCGAGGTGGAATTTGCCAGTGAGGGGAAA
P_AAX52266	1921	GTGATGGAGTATACCACCATCCTTCGGCTGCGCGAGGTGGAATTTGCCAGTGAGGGGAAA
DNA37140	1981	TATCAGTGTGTCATCTCCAATCACTTTGGTTCATCCTACTCTGTCAAAGCCAAGCTTACA
P_AAX52266	1981	TATCAGTGTGTCATCTCCAATCACTTTGGTTCATCCTACTCTGTCAAAGCCAAGCTTACA
DNA37140	2041	GTAAATATGCTTCCCTCATTCACCAAGACCCCCATGGATCTCACCATCCGAGCTGGGGCC
P_AAX52266	2041	GTAAATATGCTTCCCTCATTCACCAAGACCCCCCATGGATCTCACCATCCGAGCTGGGGCC
DNA37140	2101	ATGGCACGCTTGGAGTGTGCTGCTGGGGGCACCCAGCCCCCCAGATAGCCTGGCAGAAG
P_AAX52266	2101	ATGGCACGCTTGGAGTGTGCTGTGGGGCACCCAGCCCCCAGATAGCCTGGCAGAAG
DNA37140	2161	GATGGGGCACAGACTTCCCAGCTGCACGGGAGAGACGCATGCAT
P_AAX52266	2161	GATGGGGGCACAGACTTCCCAGCTGCACGGGAGAGACGCATGCAT
DNA37140	2221	GACGTGTTCTTTATCGTGGATGTGAAGATAGAGGACATTGGGGTATACAGCTGCACAGCT
P_AAX52266	2221	GACGTGTTCTTTATCGTGGATGTGAAGATAGAGGACATTGGGGTATACAGCTGCACAGCT
DNA37140	2281	CAGAACAGTGCAGGAAGTATTTCAGCAAATGCAACTCTGACTGTCCTAGAAACACCATCA
P_AAX52266	2281	CAGAACAGTGCAGGAAGTATTTCAGCAAATGCAACTCTGACTGTCCTAGAAACACCATCA
DNA37140	2341	TTTTTGCGGCCACTGTTGGACCGAACTGTAACCAAGGGAGAAACAGCCGTCCTACAGTGC
P_AAX52266	2341	TTTTTGCGGCCACTGTTGGACCGAACTGTAACCAAGGGAGAAACAGCCGTCCTACAGTGC
DNA37140	2401	ATTGCTGGAGGAAGCCCTCCCCTAAACTGAACTGGACCAAAGATGATAGCCCATTGGTG
P_AAX52266	2401	ATTGCTGGAGGAAGCCCTCCCCCTAAACTGAACTGGACCAAAGATGATAGCCCATTGGTG
DNA37140	2461	GTAACCGAGAGGCACTTTTTTGCAGCAGGCAATCAGCTTCTGATTATTGTGGACTCAGAT ***********************************

P_AAX52266	2461	GTAACCGAGAGGCACTTTTTTGCAGCAGGCAATCAGCTTCTGATTATTGTGGACTCAGAT
DNA37140	2521	GTCAGTGATGCTGGGAAATACACATGTGAGATGTCTAACACCCTTGGCACTGAGAGAGGA *****************************
P_AAX52266	2521	GTCAGTGATGCTGGGAAATACACATGTGAGATGTCTAACACCCTTGGCACTGAGAGAGGA
DNA37140	2581	AACGTGCGCCTCAGTGTGATCCCCACTCCAACCTGCGACTCCCCTCAGATGACAGCCCCA *****************************
P_AAX52266	2581	AACGTGCGCCTCAGTGTGATCCCCACTCCAACCTGCGACTCCCCTCAGATGACAGCCCCA
DNA37140	2641	TCGTTAGACGATGACGGATGGGCCACTGTGGGTGTCGTGATCATAGCCGTGGTTTGCTGT ************************
P_AAX52266	2641	TCGTTAGACGATGACGGATGGGCCACTGTGGGTGTCGTGATCATAGCCGTGGTTTGCTGT
DNA37140	2701	GTGGTGGGCACGTCACTCGTGTGGGTGGTCATCATATACCACACAAGGCGGAGGAATGAA ****************************
P_AAX52266	2701	GTGGTGGCACGTCACTCGTGTGGGTGGTCATCATATACCACACAAGGCGGAGGAATGAA
DNA37140	2761	GATTGCAGCATTACCAACACAGATGAGACCAACTTGCCAGCAGATATTCCTAGTTATTTG *****************************
P_AAX52266	2761	GATTGCAGCATTACCAACACAGATGAGACCAACTTGCCAGCAGATATTCCTAGTTATTTG
DNA37140	2821	TCATCTCAGGGAACGTTAGCTGACAGGCAGGATGGGTACGTGTCTTCAGAAAGTGGAAGC
P_AAX52266	2821	TCATCTCAGGGAACGTTAGCTGACAGGCAGGATGGGTACGTGTCTTCAGAAAGTGGAAGC
DNA37140	2881	CACCACCAGTTTGTCACATCTTCAGGTGCTGGATTTTTCTTACCACAACATGACAGTAGT *********************************
P_AAX52266	2881	CACCACCAGTTTGTCACATCTTCAGGTGCTGGATTTTTCTTACCACAACATGACAGTAGT
DNA37140	2941	GGGACCTGCCATATTGACAATAGCAGTGAAGCTGATGTGGAAGCTGCCACAGATCTGTTC ********************************
P_AAX52266	2941	GGGACCTGCCATATTGACAATAGCAGTGAAGCTGATGTGGAAGCTGCCACAGATCTGTTC
DNA37140	3001	CTTTGTCCGTTTTTGGGATCCACAGGCCCTATGTATTTGAAGGGAAATGTGTATGGCTCA ***********************************
P_AAX52266	3001	CTTTGTCCGTTTTTGGGATCCACAGGCCCTATGTATTTGAAGGGAAATGTGTATGGCTCA
DNA37140	3061	GATCCTTTTGAAACATATCATACAGGTTGCAGTCCTGACCCAAGAACAGTTTTAATGGAC ***********************************
P_AAX52266	3061	GATCCTTTTGAAACATATCATACAGGTTGCAGTCCTGACCCAAGAACAGTTTTAATGGAC
DNA37140	3121	CACTATGAGCCCAGTTACATAAAGAAAAAGGAGTGCTACCCATGTTCTCATCCTTCAGAA **********************************
P_AAX52266.	3121	CACTATGAGCCCAGTTACATAAAGAAAAAGGAGTGCTACCCATGTTCTCATCCTTCAGAA
DNA37140		GAATCCTGCGAACGGAGCTTCAGTAATATATCGTGGCCTTCACATGTGAGGAAGCTACTT *********************************
P_AAX52266		GAATCCTGCGAACGGAGCTTCAGTAATATATCGTGGCCTTCACATGTGAGGAAGCTACTT
DNA37140		AACACTAGTTACTCTCACAATGAAGGACCTGGAATGAAAAATCTGTGTCTAAACAAGTCC ***********************************
P_AAX52266		AACACTAGTTACTCTCACAATGAAGGACCTGGAATGAAAAATCTGTGTCTAAACAAGTCC
DNA37140		TCTTTAGATTTTAGTGCAAATCCAGAGCCAGCGTCGGTTGCCTCGAGTAATTCTTTCATG ************************************
P_AAX52266	3301	TCTTTAGATTTTAGTGCAAATCCAGAGCCAGCGTCGGTTGCCTCGAGTAATTCTTTCATG

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3361 GGTACCTTTGGAAAAGCTCTCAGGAGACCTCACCTAGATGCCTATTCAAGCTTTGGACAG
  DNA37140
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           3361 GGTACCTTTGGAAAAGCTCTCAGGAGACCTCACCTAGATGCCTATTCAAGCTTTGGACAG
 P AAX52266
  DNA37140
           3421 CCATCAGATTGTCAGCCAAGAGCCTTTTATTTGAAAGCTCATTCTTCCCCAGACTTGGAC
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P AAX52266
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P AAX52266
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  DNA37140
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              ******************
P AAX52266
          3841 TATCTTACGTAAATTAATGATATAAATCATGATTATTTTATGTATTTTATAATGCCAGA
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P AAX52266
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 DNA37140
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              ******************
P AAX52266
          3961 TTTTAAATAGAAGTTACTTCATTATATTTTGCACATTATATTTAATAAAATGTGTCAATT
 DNA37140
         4021 TGAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAA
              *****
P AAX52266
         4021 TGAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAAA
>4 AX098387 Sequence 13 from Patent W00119991. (4053 bp) [1 seg]
Score = 4053 (8034 bits), Expect = 0.0
Identities = 4053/4053 (100%), at 1,1-4053,4053, Strand +/+
 DNA37140
            1 AGCCGACGCTGCTCAAGCTGCAACTCTGTTGCAGTTGGCAGTTCTTTTCGGTTTCCCTCC
              **********************
 AX098387
            1 \ \ \mathsf{AGCCGACGCTGCTCAAGCTGCAACTCTGTTGCAGTTGGCAGTTCTTTTCGGTTTCCCTCC}
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DNA37140	61	TGCTGTTTGGGGGCATGAAAGGGCTTCGCCGCCGGGAGTAAAAGAAGAATTGACCGGC
AX098387	61	TGCTGTTTGGGGGCATGAAAGGGCTTCGCCGCCGGGAGTAAAAGAAGAATTGACCGGGC
DNA37140	121	AGCGCGAGGGAGGGCGCGCGACGCGACGCGAGGGCGGGC
AX098387	121	AGCGCGAGGGAGGGCGCGCGCGACGCGAGGGCGGGCGTGCACCCTCGGCTGGAAGT
DNA37140	181	TTGTGCCGGGCCCGAGCGCGCCGGCTGGGAGCTTCGGGTAGAGACCTAGGCCGCTGG ******************************
AX098387	181	TTGTGCCGGGCCCCGAGCGCCGCCGGCTGGGAGCTTCGGGTAGAGACCTAGGCCGCTGG
DNA37140	241	ACCGCGATGAGCGCCGAGCCTCCGTGCGCGCGCGCGGGGTTGGGGCTGCTGTGC **********
AX098387	241	ACCGCGATGAGCGCCGAGCCTCCGTGCGCGCGCGCGGGGTTGGGGCTGCTGTGC
DNA37140		GCGGTGCTGGGCGCCGGTCCGACAGCGGCGGTCGCGGGGAACTCGGGCAGCCC *****************************
AX098387		GCGGTGCTGGGCGCCTGGCCGGTCCGACAGCGGCGGTCGCGGGGAACTCGGGCAGCCC
DNA37140	361	TCTGGGGTAGCCGCCGAGCGCCCATGCCCCACTACCTGCCGCTGCCTCGGGGACCTGCTG ********************************
AX098387		TCTGGGGTAGCCGCCGAGCGCCCATGCCCACTACCTGCCGCTGCCTCGGGGACCTGCTG
DNA37140		GACTGCAGTCGTAAGCGGCTAGCGCGTCTTCCCGAGCCACTCCCGTCCTGGGTCGCTCGG ******************
AX098387 DNA37140		GACTGCAGTCGTAAGCGGCTAGCGCGTCTTCCCGAGCCACTCCCGTCCTGGGTCGCTCGG
AX098387		CTGGACTTAAGTCACAACAGATTATCTTTCATCAAGGCAAGTTCCATGAGCCACCTTCAA *****************************
DNA37140		CTGGACTTAAGTCACAACAGATTATCTTTCATCAAGGCAAGTTCCATGAGCCACCTTCAA AGCCTTCGAGAAGTGAAACTGAACAACAATGAATTGGAGACCATTCCAAATCTGGGACCA
		**************
AX098387		AGCCTTCGAGAAGTGAAACTGAACAACAATGAATTGGAGACCATTCCAAATCTGGGACCA
DNA37140		GTCTCGGCAAATATTACACTTCTCTCTTGGCTGGAAACAGGATTGTTGAAATACTCCCT ****************************
AX098387		GTCTCGGCAAATATTACACTTCTCCTTGGCTGGAAACAGGATTGTTGAAATACTCCCT
DNA37140		GAACATCTGAAAGAGTTTCAGTCCCTTGAAACTTTGGACCTTAGCAGCAACAATATTTCA **************************
AX098387 DNA37140		GAACATCTGAAAGAGTTTCAGTCCCTTGAAACTTTTGGACCTTAGCAGCAACAATATTTCA
		GAGCTCCAAACTGCATTTCCAGCCCTACAGCTCAAATATCTGTATCTCAACAGCAACCGA **************************
AX098387	721	GAGCTCCAAACTGCATTTCCAGCCCTACAGCTCAAATATCTGTATCTCAACAGCAACCGA
DNA37140	781	GTCACATCAATGGAACCTGGGTATTTTGACAATTTGGCCAACACACTCCTTGTGTTAAAG ****************************
AX098387	781	GTCACATCAATGGAACCTGGGTATTTTGACAATTTGGCCAACACACTCCTTGTGTTAAAG
DNA37140	841	CTGAACAGGAACCGAATCTCAGCTATCCCACCCAAGATGTTTAAACTGCCCCAACTGCAA
AX098387	841	CTGAACAGGAACCGAATCTCAGCTATCCCACCCAAGATGTTTAAACTGCCCCAACTGCAA
DNA37140	901	CATCTCGAATTGAACCGAAACAAGATTAAAAATGTAGATGGACTGACATTCCAAGGCCTT

		******************
AX098387	901	CATCTCGAATTGAACCGAAACAAGATTAAAAATGTAGATGGACTGACATTCCAAGGCCTT
DNA37140	961	GGTGCTCTGAAGTCTCTGAAAATGCAAAGAAATGGAGTAACGAAACTTATGGATGG
AX098387	961	GGTGCTCTGAAGTCTCTGAAAATGCAAAGAAATGGAGTAACGAAACTTATGGATGG
DNA37140	1021	TTTTGGGGGCTGAGCAACATGGAAATTTTGCAGCTGGACCATAACAACCTAACAGAGATT
AX098387	1021	TTTTGGGGGCTGAGCAACATGGAAATTTTGCAGCTGGACCATAACAACCTAACAGAGATT
DNA37140	1081	ACCAAAGGCTGGCTTTACGGCTTGCTGATGCTGCAGGAACTTCATCTCAGCCAAAATGCC
AX098387	1081	ACCAAAGGCTGGCTTTACGGCTTGCTGATGCTGCAGGAACTTCATCTCAGCCAAAATGCC
DNA37140	1141	ATCAACAGGATCAGCCCTGATGCCTGGGAGTTCTGCCAGAAGCTCAGTGAGCTGGACCTA
AX098387	1141	ATCAACAGGATCAGCCCTGATGCCTGGGAGTTCTGCCAGAAGCTCAGTGAGCTGGACCTA
DNA37140	1201	ACTTTCAATCACTTATCAAGGTTAGATGATTCAAGCTTCCTTGGCCTAAGCTTACTAAAT
AX098387	1201	ACTTTCAATCACTTATCAAGGTTAGATGATTCAAGCTTCCTTGGCCTAAGCTTACTAAAT
DNA37140	1261	ACACTGCACATTGGGAACAACAGAGTCAGCTACATTGCTGATTGTGCCTTCCGGGGGCTT ********************
AX098387	1261	ACACTGCACATTGGGAACAACAGAGTCAGCTACATTGCTGATTGTGCCTTCCGGGGGCTT
DNA37140	1321	TCCAGTTTAAAGACTTTGGATCTGAAGAACAATGAAATTTCCTGGACTATTGAAGACATG
AX098387	1321	TCCAGTTTAAAGACTTTGGATCTGAAGAACAATGAAATTTCCTGGACTATTGAAGACATG
DNA37140	1381	AATGGTGCTTTCTCTGGGCTTGACAAACTGAGGCGACTGATACTCCAAGGAAATCGGATC ************************************
AX098387	1381	AATGGTGCTTTCTCTGGGCTTGACAAACTGAGGCGACTGATACTCCAAGGAAATCGGATC
DNA37140	1441	CGTTCTATTACTAAAAAAGCCTTCACTGGTTTGGATGCATTGGAGCATCTAGACCTGAGT **********************************
AX098387	1441	CGTTCTATTACTAAAAAAGCCTTCACTGGTTTGGATGCATTGGAGCATCTAGACCTGAGT
DNA37140	1501	GACAACGCAATCATGTCTTTACAAGGCAATGCATTTTCACAAATGAAGAAACTGCAACAA
AX098387	1501	GACAACGCAATCATGTCTTTACAAGGCAATGCATTTTCACAAATGAAGAAACTGCAACAA
DNA37140	1561	TTGCATTTAAATACATCAAGCCTTTTGTGCGATTGCCAGCTAAAATGGCTCCCACAGTGG **********************************
AX098387	1561	TTGCATTTAAATACATCAAGCCTTTTGTGCGATTGCCAGCTAAAATGGCTCCCACAGTGG
DNA37140	1621	GTGGCGGAAAACAACTTTCAGAGCTTTGTAAATGCCAGTTGTGCCCATCCTCAGCTGCTA ************************************
AX098387		GTGGCGGAAAACAACTTTCAGAGCTTTGTAAATGCCAGTTGTGCCCATCCTCAGCTGCTA
DNA37140		AAAGGAAGAAGCATTTTTGCTGTTAGCCCAGATGGCTTTGTGTGTG
AX098387		AAAGGAAGAAGCATTTTTGCTGTTAGCCCAGATGGCTTTGTGTGTG
DNA37140	1741	CCCCAGATCACGGTTCAGCCAGAAACACAGTCGGCAATAAAAGGTTCCAATTTGAGTTTC **********************************

AX098387	1741	CCCCAGATCACGGTTCAGCCAGAAACACAGTCGGCAATAAAAGGTTCCAATTTGAGTTTC
DNA37140	1801	ATCTGCTCAGCTGCCAGCAGCAGTGATTCCCCCAATGACTTTTGCTTGGAAAAAAGACAAT
AX098387	1801	${\tt ATCTGCTCAGCTGCCAGCAGCAGTGATTCCCCCAATGACTTTTGCTTGGAAAAAAAGACAAT}$
DNA37140	1861	GAACTACTGCATGATGCTGAAATGGAAAATTATGCACACCTCCGGGCCCAAGGTGGCGAG
AX098387	1861	GAACTACTGCATGATGCTGAAATGGAAAATTATGCACACCTCCGGGCCCAAGGTGGCGAG
DNA37140	1921	GTGATGGAGTATACCACCATCCTTCGGCTGCGCGAGGTGGAATTTGCCAGTGAGGGGAAA ***************************
AX098387	1921	GTGATGGAGTATACCACCATCCTTCGGCTGCGCGAGGTGGAATTTGCCAGTGAGGGGAAA
DNA37140	1981	TATCAGTGTGTCATCTCCAATCACTTTGGTTCATCCTACTCTGTCAAAGCCAAGCTTACA
AX098387	1981	TATCAGTGTGTCATCTCCAATCACTTTGGTTCATCCTACTCTGTCAAAGCCAAGCTTACA
DNA37140	2041	GTAAATATGCTTCCCTCATTCACCAAGACCCCCATGGATCTCACCATCCGAGCTGGGGCC ******************************
AX098387	2041	GTAAATATGCTTCCCTCATTCACCAAGACCCCCATGGATCTCACCATCCGAGCTGGGGCC
DNA37140	2101	ATGGCACGCTTGGAGTGTGCTGCTGTGGGGCACCCAGCCCCCAGATAGCCTGGCAGAAG
AX098387	2101	$\tt ATGGCACGCTTGGAGTGTGCTGCTGTGGGGCACCCAGCCCCCCAGATAGCCTGGCAGAAG$
DNA37140	2161	GATGGGGGCACAGACTTCCCAGCTGCACGGGAGAGACGCATGCAT
AX098387	2161	GATGGGGGCACAGACTTCCCAGCTGCACGGGAGAGACGCATGCAT
DNA37140	2221	GACGTGTTCTTTATCGTGGATGTGAAGATAGAGGACATTGGGGTATACAGCTGCACAGCT
AX098387	2221	GACGTGTTCTTTATCGTGGATGTGAAGATAGAGGACATTGGGGTATACAGCTGCACAGCT
DNA37140	2281	CAGAACAGTGCAGGAAGTATTTCAGCAAATGCAACTCTGACTGTCCTAGAAACACCATCA
AX098387	2281	CAGAACAGTGCAGGAAGTATTTCAGCAAATGCAACTCTGACTGTCCTAGAAACACCATCA
DNA37140	2341	TTTTTGCGGCCACTGTTGGACCGAACTGTAACCAAGGGAGAAACAGCCGTCCTACAGTGC ***********************************
AX098387	2341	$\tt TTTTTGCGGCCACTGTTGGACCGAACTGTAACCAAGGGAGAAACAGCCGTCCTACAGTGC$
DNA37140	2401	ATTGCTGGAGGAAGCCCTCCCCCTAAACTGAACTGGACCAAAGATGATAGCCCATTGGTG
AX098387	2401	ATTGCTGGAGGAAGCCCTCCCCCTAAACTGAACTGGACCAAAGATGATAGCCCATTGGTG
DNA37140	2461	GTAACCGAGAGGCACTTTTTTGCAGCAGGCAATCAGCTTCTGATTATTGTGGACTCAGAT
AX098387	2461	GTAACCGAGAGGCACTTTTTTGCAGCAGGCAATCAGCTTCTGATTATTGTGGACTCAGAT
DNA37140	2521	GTCAGTGATGCTGGGAAATACACATGTGAGATGTCTAACACCCTTGGCACTGAGAGAGA
AX098387	2521	GTCAGTGATGCTGGGAAATACACATGTGAGATGTCTAACACCCTTGGCACTGAGAGAGGA
DNA37140	2581	AACGTGCGCCTCAGTGTGATCCCCACTCCAACCTGCGACTCCCCTCAGATGACAGCCCCA
AX098387	2581	AACGTGCGCCTCAGTGTGATCCCCACTCCAACCTGCGACTCCCCTCAGATGACAGCCCCA

DNA37140	2641	TCGTTAGACGATGACGGATGGGCCACTGTGGGTGTCGTGATCATAGCCGTGGTTTGCTGT
AX098387	2641	TCGTTAGACGATGACGGATGGGCCACTGTGGGTGTCGTGATCATAGCCGTGGTTTGCTGT
DNA37140	2701	GTGGTGGCACGTCACTCGTGTGGGTGGTCATCATATACCACACAAGGCGGAGGAATGAA
AX098387	2701	GTGGTGGCACGTCACTCGTGTGGGTGGTCATCATATACCACACAAGGCGGAGGAATGAA
DNA37140	2761	GATTGCAGCATTACCAACACAGATGAGACCAACTTGCCAGCAGATATTCCTAGTTATTTG
AX098387	2761	GATTGCAGCATTACCAACACAGATGAGACCAACTTGCCAGCAGATATTCCTAGTTATTTG
DNA37140	2821	TCATCTCAGGGAACGTTAGCTGACAGGCAGGATGGGTACGTGTCTTCAGAAAGTGGAAGC *********************************
AX098387	2821	TCATCTCAGGGAACGTTAGCTGACAGGCAGGATGGGTACGTGTCTTCAGAAAGTGGAAGC
DNA37140	2881	CACCACCAGTTTGTCACATCTTCAGGTGCTGGATTTTTCTTACCACAACATGACAGTAGT
AX098387	2881	CACCACCAGTTTGTCACATCTTCAGGTGCTGGATTTTTCTTACCACAACATGACAGTAGT
DNA37140	2941	GGGACCTGCCATATTGACAATAGCAGTGAAGCTGATGTGGAAGCTGCCACAGATCTGTTC ********************************
AX098387	2941	GGGACCTGCCATATTGACAATAGCAGTGAAGCTGATGTGGAAGCTGCCACAGATCTGTTC
DNA37140	3001	CTTTGTCCGTTTTTGGGATCCACAGGCCCTATGTATTTGAAGGGAAATGTGTATGGCTCA
AX098387	3001	CTTTGTCCGTTTTTGGGATCCACAGGCCCTATGTATTTGAAGGGAAATGTGTATGGCTCA
DNA37140	3061	GATCCTTTTGAAACATATCATACAGGTTGCAGTCCTGACCCAAGAACAGTTTTAATGGAC ***********************************
AX098387	3061	GATCCTTTTGAAACATATCATACAGGTTGCAGTCCTGACCCAAGAACAGTTTTAATGGAC
DNA37140	3121	CACTATGAGCCCAGTTACATAAAGAAAAAGGAGTGCTACCCATGTTCTCATCCTTCAGAA
AX098387	3121	CACTATGAGCCCAGTTACATAAAGAAAAAGGAGTGCTACCCATGTTCTCATCCTTCAGAA
DNA37140	3181	GAATCCTGCGAACGGAGCTTCAGTAATATATCGTGGCCTTCACATGTGAGGAAGCTACTT
AX098387	3181	GAATCCTGCGAACGGAGCTTCAGTAATATCGTGGCCTTCACATGTGAGGAAGCTACTT
DNA37140	3241	AACACTAGTTACTCTCACAATGAAGGACCTGGAATGAAAAATCTGTGTCTAAACAAGTCC ***********************************
AX098387	3241	AACACTAGTTACTCTCACAATGAAGGACCTGGAATGAAAAATCTGTGTCTAAACAAGTCC
DNA37140	3301	TCTTTAGATTTTAGTGCAAATCCAGAGCCAGCGTCGGTTGCCTCGAGTAATTCTTTCATG ************************************
AX098387	3301	TCTTTAGATTTTAGTGCAAATCCAGAGCCAGCGTCGGTTGCCTCGAGTAATTCTTTCATG
DNA37140	3361	GGTACCTTTGGAAAAGCTCTCAGGAGACCTCACCTAGATGCCTATTCAAGCTTTGGACAG *********************************
AX098387	3361	GGTACCTTTGGAAAAGCTCTCAGGAGACCTCACCTAGATGCCTATTCAAGCTTTGGACAG
DNA37140	3421	CCATCAGATTGTCAGCCAAGAGCCTTTTATTTGAAAGCTCATTCTTCCCCAGACTTGGAC
AX098387	3421	CCATCAGATTGTCAGCCAAGAGCCTTTTATTTGAAAGCTCATTCTTCCCCAGACTTGGAC